Abstract

Ordinal data are extremely used in psychology. Common examples include Likert items or variables where categories are ordered without assuming a common distance between labels. Despite the prevalence of ordinal data, a majority of published psychological research relies on applying metric models (e.g., standard linear regression) even when the dependent variable is ordinal. This approach leads to several statistical drawbacks. Similar to what happens with binary data (e.g., accuracies in a cognitive experiment), using metric models instead of e.g. logistic regression is probably caused by several reasons. Firstly, parameter interpretation can be challenging. Ordinal models commonly used a link function (e.g., logit) for model fitting thus transforming the parameters scale. This is even more critical when estimating models with multiple parameters or interactions. Secondly, nowadays journals require power or sensitivity analysis when planning or evaluating a research work. For common statistical models such as t-tests or the analysis of variance, there are tools to easily estimate the statistical power. For ordinal regressions, estimating the statistical power often requires simulating data thus clearly understanding the statistical background and implementing a Monte Carlo simulation. The objective of this tutorial is to present the essential statistical background of ordinal regression, highlighting the research areas that would benefit more by introducing this model. In particular, we present how to interpret the model parameters with categorical and numerical variables also linking them to common standardized effect size measures. Then we propose methods to report the results, especially focusing on graphical representation. Finally, we present how to estimate the statistical power using Monte Carlo simulations with suggestions on setting parameters in a meaningful way. Through the tutorial, we use the R programming language using specific packages for ordinal models and sharing also ad-hoc developed functions supporting each step. For the sake of reproducibility and to empower researchers in analyzing their studies autonomously, all data and scripts used in the article have been made publicly available.

Keywords: ordinal, simulations, power

Word count: X

Ordinal regression models made easy. A tutorial on parameter interpretation, data simulation, and power analysis.

Introduction

Psychological research make an extensive use of ordinal data. One of the main reason is probably the usage of Likert scales (Likert, 1932). Ordinal data, as defined by Stevens (1946), belongs to a specific type of measurement scale where ordered numbers are assigned to a variable. Beyond Likert-like items, in Psychology there are several applications of ordinal scales. For instance, sociodemographic variables like educational levels or socioeconomic status, as well as general ratings such as pain severity, agreement with a statement, or the evaluation of sensory experiences.

In contrast to nominal scales, the labels in ordinal scales are ordered. Unlike interval or ratio scales, there is no explicit assumption about the distance between labels. An example is asking people the degree of agreement about a certain statement using a scale from 1 (no agreement) to 7 (total agreement). Answering 4 compared to 2 suggest an higher agreement but we cannot affirm that there is two times the agreement compared to the second answer. Stevens (1946) and Kemp and Grace (2021) suggested that for ordinal variables is appropriate to calculate ranks-based descriptive statistics (e.g., median or percentiles) instead of metric statistics (e.g., mean or standard deviation) and using appropriate inferential tools (Agresti, 2010; e.g., N. Cliff, 1996). This distinction in terms of the appropriateness of certain descriptive statistics is also relevant when modeling data. Treating ordinal data as metric refers to assuming the labels as actual integer numbers thus assuming a fixed and know distance between levels (Liddell & Kruschke, 2018). More generally, Norman Cliff (2016) suggest that most of the research questions in behavioral sciences can be considered as ordinal (is the score x higher than the score y?) concerning variables where the most appropriate measurement scale is probably ordinal.

In Psychology especially when using item-based measures (questionnaires, surveys,

etc.) the common practice is using a normal linear regression that makes an explicit assumption about metric features of the response variable. Liddell and Kruschke (2018) reviewed the psychological literature using likert-based measures and reported how the majority of papers used metric-based statistical models. In the same work, Liddell and Kruschke (2018) showed extensive examples and simulations about the potential pitfalls of treating an ordinal variable as metric (but see Robitzsch, 2020 for an alternative perspective). They reported problems in terms of lack of power, inversion of the effects (e.g., finding a negative effect when the true effect is positive) and distorted effect size estimates. Some authors suggested that individual ordinal items and a collection of ordinal (averaged or summed) items can safely be considered and analyzed as metric (Carifio & Perla, 2008; Carifio & Perla, 2007; but see Jamieson, 2004). Despite Liddell and Kruschke (2018) provide some example that averaging ordinal items and applying metric models is not appropriate, in the current paper we discuss only cases where there is a single ordinal outcome (e.g., a single item, question, etc.).

For the tutorial, we assume that the reader is already familiar with basic R programming and introductory theory about linear regression. Despite the practical and applicative spirit of the tutorial, we still introduced some statistical theory and equations supported by translation into R code. Similar to other tutorials (DeBruine & Barr, 2021; Gambarota & Altoè, 2023), we believe this is the best educational approach especially with the involvement of Monte Carlo simulations.

In the first part we introduce the ordinal models explaining the general structure and assumptions. Then we move with model fitting and parameters interpretation. Finally we introduce the simulation approach for common research scenarios and an application to power analysis. We used R [Version 4.3.1; R Core Team (2023)] for the code, figures and tables. Details about R packages an The tutorial is supported by a set of custom R functions available on Github. Other examples and model extensions along with details about used R packages are available on the Github repository (supplementary materials).

For the tutorial we used the following packages.

Ordinal regression models

While Liddell and Kruschke (2018) proposed a specific modeling approach, there exists a class of regression models that consider the ordinal nature of the response variable without relying on metric assumptions. We can name this general class of models as ordinal regression. The actual statistical nomenclature can be confusing mainly because there are several types of models with different assumptions and structures (Tutz, 2022). Tutz (2022) and Bürkner and Vuorre (2019) provide a clear and updated taxonomy of ordinal regression models.

We can identify three main classes of models: cumulative models (Agresti, 2010; CM, McCullagh, 1980), sequential models (Tutz, 1990), and adjacent category models. Among these, the cumulative model is the most widely used, assuming the existence of a latent variable. This latent variable, categorized using a set of thresholds, produces the observed ordinal variable. The sequential model as suggested by the name is appropriate when modeling sequential processes. Assuming to have five response options, the sequential model assume that responding "3" assume a sequential process where steps "1" and "2" are already reached. A clear example is proposed by Bürkner and Vuorre (2019) where the marriage duration in years is predicted as a function of some explanatory variables. For each level of the response variable there is a latent distribution where the step between a marriage year k = 1 and the next years k > 1 is modeled by the sequential model. When comparing k with k > 1, everything lower than k is assumed to be already reached (Tutz & Berger, 2020).

The adjacent category model compare the category k with k+1 still assuming a latent distribution for each k. As suggested by Tutz (2022) the adjacent-category model can be seen as a series of binary binomial regressions taking into account the order of the

categories. Bürkner and Vuorre (2019) suggested that adjacent-category model can be chosen for its mathematical convenience and there is no a clear empirical interpretation as for the cumulative vs sequential model.

In the current paper we put the focus on the CM for several reasons. The first reason is that the latent formulation of the model is particularly convenient both for parameter interpretation and data simulation. The second reason is that several psychological variables can be formalized as a latent continuous variable observed as an ordinal item. Furthermore, CM are also used to model data under a signal detection theory framework (e.g., DeCarlo, 2010). Figure 1 depicts the overall structure of the *cumulative* model.

Model notation

In this section we introduce some notation for the CM that is used through the paper. We proposed a notation as consistent as possible with the literature and the ordinal package (Christensen, 2022). We define Y_k as the observed ordinal variable with k levels and Y^* is the underlying latent variable. The latent variable is segmented using k-1 thresholds $\alpha_1, \ldots, \alpha_{k-1}$. Similarly to the generalized linear models framework (e.g., Fox, 2015), we define $g(x) = \eta$ as the link function that maps probabilities into the linear predictor η . To transform back η into probabilities we use the inverse of the link function $x = g^{-1}(\eta)$. The specific link function define the type of model and require a different R function. For example, when assuming a Gaussian distribution, fitting a model with a probit link function entails using the cumulative distribution function $g(x) = \Phi^{-1}(x) = \eta$ and the inverse of the link function is the inverse cumulative distribution function (or quantile function) defined $x = g^{-1}(\eta) = \Phi(\eta)$. When modelling an ordinal variable in a cumulative link model we actually modelling the cumulative probability $P(Y \le k), k = 1, \ldots, k-1$. Equation (1) shows the general cumulative model including

¹ As done by Agresti (2010), when referring to $P(Y \le k)$ we are implicitly conditioning on a particular **X** value $P(Y \le k | \mathbf{X}_i)$

predictors \mathbf{X} and regression coefficients β . The minus sign in $\mathbf{X}\beta$ is used to interpret the β_j as in the standard regression models where higher β values corresponds to increased probability of responding higher k categories (Agresti, 2010).

$$P(Y \le k) = g^{-1}(\alpha_k - \mathbf{X}\beta) \quad k = 1, \dots, k - 1 \tag{1}$$

The $X\beta$ is the linear predictor η that is the cumulative probability $P(Y \leq k)$ transformed using the link function $g(\cdot)$. To obtain the probability of a single outcome P(Y = k) we can compute the difference between cumulative probabilities as shown in Equation (2).

$$P(Y=k) = g^{-1}(\alpha_k - \eta) - g^{-1}(\alpha_{k-1} - \eta), \quad k = 1, \dots, k-1$$
 (2)

There are always two special cases when computing the probability of a single outcome Y that is when Y=1 and Y=k. In the first case the cumulative probability is calculated from $-\infty$ that is the same as temporary assuming an $\alpha_0=-\infty$. In the second case (Y=1) the probability is calculated as $P(Y=k)=1-g^{-1}(\alpha_{k-1}-\eta)$ that is the same as assuming a temporary threshold $\alpha_k=+\infty$. The Figure 1 shows how the single probabilities of the ordinal outcome are calculated from cumulative probabilities.

[Figure 1 about here]

The same model can be written in the so-called latent formulation as reported in Equation (3). The model is no longer directly about the cumulative probabilities but focused on the continuous latent variable Y^* as a function of the linear predictor $\eta = \mathbf{X}\beta$ similar to a standard linear regression.

$$\mathbf{Y}^{\star} = \mathbf{X}\beta + \tag{3}$$

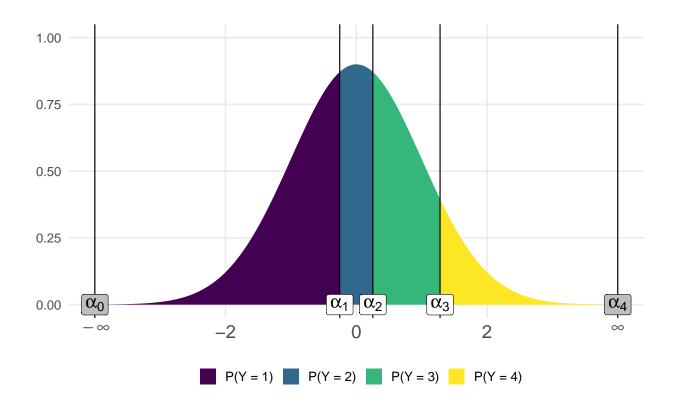


Figure 1. Relationship between cumulative probabilities and ordinal outcomes. The Y^* latent variable is segmented into k levels using k-1 thresholds (α). The distance between thresholds determine the probability of each outcome. The grey thresholds are only used to compute the Y=1 and Y=k probabilities.

The crucial part is ϵ that is the random component of the model coming from a certain probability distribution. For a *probit* model, errors are sampled from a standard Gaussian distribution while for a *logit* model from a standard logistic distribution. Following the notation by Tutz (2022), the observed ordinal value $Y_i = k$ comes from Y_i^* belonging to the interval defined by the thresholds $Y_i = k \iff \alpha_{k-1} < Y_i^* < \alpha_k$ where $-\infty = \alpha_0 < \alpha_1 < \dots < \alpha_{k-1} < \alpha_k = \infty$.

In the basic version of the model, the thresholds α_k are considered as fixed and being part of the measurement procedure (Liddell & Kruschke, 2018) and do not vary as a function of the predictors. In a more sophisticated version of the model called location-shift (Tutz, 2022), both the location μ and the thresholds α_k can vary as a function of the

predictors.

The model can be also formalized in alternative ways. Liddell and Kruschke (2018) and Kruschke (2015) proposed a Bayesian version of the model with a different threshold parametrization. Gelman, Hill, and Vehtari (2020) proposed three alternative parametrizations focusing on different definition of the thresholds.

Tutz (2022) described also another version of the model called location-scale (Cox, 1995; Rigby & Stasinopoulos, 2005; Tutz, 2022) where the location μ and the scale σ^2 of the distribution can vary as a function of the predictors.

Link function

The cumulative link model implemented in Equations (1) and (3) can be considered the general formulation that requires specifying the link function $g(\cdot)$ or the errors distribution $\epsilon_i \sim D(\mu, \sigma^2)$. Among several available functions the logit and probit models are the most common. The logit model use a logit link function and a logistic distribution as latent variable. On the other side, the probit model assume a latent Gaussian distribution using the inverse of the cumulative distribution (Φ^{-1}) function as link function.

The two models provide similar results with a different parameters interpretation. In the next sections we will illustrate the differences and simulation strategies. Figure 2 depicts the two distributions while Table L'output è 1 summarise the presented cumulative models with the proposed link function and the corresponding R code.

In terms of parameters, both distributions can be defined with a location μ and a scale s parameter. The standard normal distribution has $\mu=0$ and s=1. Furthermore the variance corresponds to the scale $s^2=\sigma^2=1$. The variance of the logistic distribution is $\sigma^2=\frac{s^2\pi^2}{3}$. The standard logistic distribution has $\mu=0$ and $s^2=1$ thus the standard deviation simplified to $\frac{\pi}{\sqrt{3}}\approx 1.81$. In practical terms, fixing μ and s lead to an higher standard deviation for the logistic distribution.

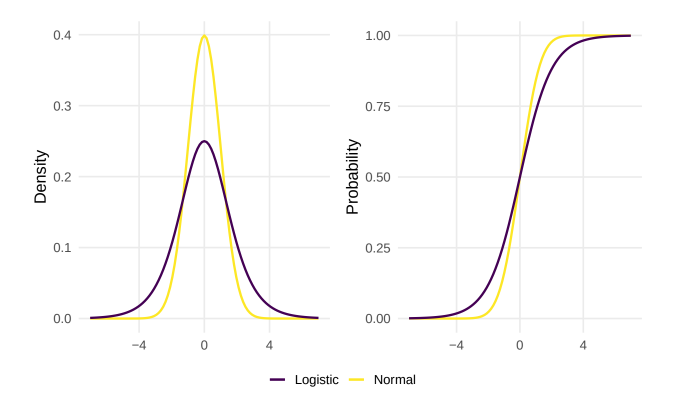


Figure 2. Difference between *logit* and *probit* models. On the left the probability density function (PDF). On the right the cumulative distribution function (CDF).

[Figure 2 about here]

Model fitting

For fitting the cumulative models we used the ordinal package (Christensen, 2019). Despite the presence of other possibilities the ordinal package provide the most complete and intuitive way to implement the ordinal models. The syntax is very similar to standard linear models in R and default functions to calculate predictions, perform model comparison, extract relevant model information are implemented similarly to standard regression modelling.²

² For a very complete overview of the ordinal package see Christensen (2019) and the Package documentation https://cran.r-project.org/web/packages/ordinal/ordinal.pdf

Table 1
(ref:tab-model-summary-docx)

	Link Func	tion $g(.)$	Inverse Link Function $g^{-1}(.)$		
Model	Equation	R Code	Equation	R Code	
Cumulative Probit	$z = \Phi^{-1}(p)$	qnorm()	$p=\Phi(z)$	pnorm()	

The function to fit the model is clm() and the model equation is specified using the R formula syntax y ~ x where y is the ordinal dependent variable and x is one or more predictors eventually including also the interactions. The package also implements mixed-effects models (see the clmm() function) including random intercepts and slopes but the topic is beyond the scope of the current tutorial.

When fitting the model the crucial arguments are the formula, the link function and the data. More advanced arguments are the nominal, scale and threshold.

- formula: the formula y ~ x with the dependent variable and predictors.
- link: is the link function. In this tutorial we consider only the *logit* and *probit* link but other link functions are available.
- data: is the dataset with variables included in the formula
- nominal: formula with predictors where the proportional odds assumption (See Section) is relaxed (i.e., partial or non proportional odds)
- scale: formula with predictors for the scale (standard deviation) parameter. This
 argument allow to fit a scale-location model (see supplementary materials). The main
 formula argument refers to predictors on the location parameter (i.e., the mean μ).
- threshold: different structures for estimating the thresholds. The default is
 threshold = "flexible" where k 1 threshold (where k is the number of ordinal levels for Y) are estimated.

We can start by fitting a simple model, highlighting the crucial parameters where the detailed explanation will be expanded in the next sections. Table 2 contains simulated data from n=100 participants rating the agreement about a certain item with k=4 ordered options. The participants are divided into two groups $(x_a \text{ and } x_b)$. We can fit a cumulative link model with $\mathtt{clm}()$ function and check the model summary.

Table 2

(ref:tab-dataset-example-docx)

Group	Mean	Median	SD	Y 1	$\mathbf{Y2}$	Y3	Y4
a	2.16	2	1.09	19 $(p = 0.38)$	11 (p = 0.22)	13 (p = 0.26)	7 (p = 0.14)
b	3.12	4	1.1	7 (p = 0.14)	6 (p = 0.12)	11 (p = 0.22)	26 (p = 0.52)

```
fit <- clm(y ~ x, data = dat, link = "logit")</pre>
summary(fit)
\# formula: y \sim x
#> data: dat
#> link threshold nobs logLik AIC     niter max.grad cond.H
   logit flexible 100 -127.04 262.08 4(0) 5.42e-07 2.0e+01
#> Coefficients:
   Estimate Std. Error z value Pr(>|z|)
#> xb 1.5929 0.3895 4.09 4.32e-05 ***
#> Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
#>
#> Threshold coefficients:
    Estimate Std. Error z value
#> 1/2 -0.4243 0.2745 -1.546
#> 2|3 0.4293 0.2711 1.584
#> 3/4 1.5851 0.3165 5.008
```

The two main sections of the model summary are the *Coefficients* section reporting the regression coefficients β and the *Threshold* section reporting the α estimation. Given

that k=4 we have k-1=3 thresholds and one β associated with the x effect. As in standard regression models, when x is a categorical predictor with j level, we will estimate j-1 regression coefficients (plus the intercept term) where the interpretation depends on the contrast coding (see Schad, Vasishth, Hohenstein, & Kliegl, 2020). In R the default is the dummy coding where a factor of j levels is converted into j-1 dummy variables. By default, the first level of the factor is taken as the reference level and the j-1 coefficients represent the comparison between the other levels and the reference.

Interpreting parameters

Logit Model. Odds and odds ratio

To understand the logit model we need to introduce odds and odds ratio. The odds of a probability p is defined as p/(1-p) thus the probability of success divided by the probability of failure. The odds takes value ranging from 0 to ∞ . For instance, with a probability of p=0.8, we have odds of 4, indicating that there are four successes for each failure. The same as having p=0.2 and an odds of 0.25 means that for each 0.25 successes we have a failure or that we have 4 failures for each success. When comparing two groups or conditions we can compare the two odds calculating an odds ratio. The odds ratio is the mostly used statistics to compare groups or conditions on a binary outcome. An odds ratio of 4 means that the odds of success at the numerator is 4 times higher than the odds of success at the denominator. The Figure 3 shows the relationship between probabilities and odds. The logit transformation is about taking the logarithm of the odds creating a symmetric function ranging from $-\infty$ to ∞ with p=0.5 as the midpoint because $\log(0.5/(1-0.5))=0$.

[Figure 3 about here]

Odds and odds ratios are clearly defined with k=2 outcomes. With an ordinal variable and a cumulative model we can use the cumulative odds ratio. With e.g. k=4

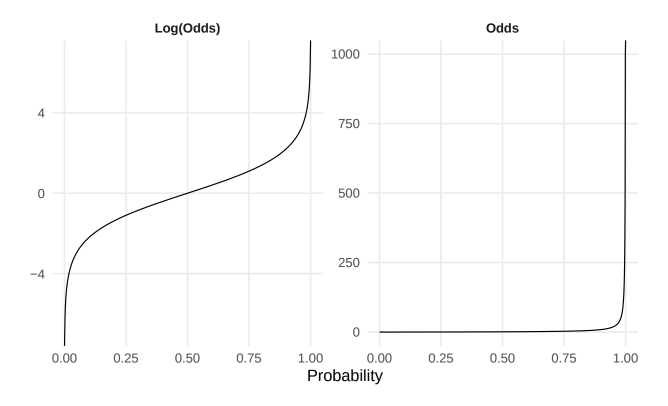


Figure 3. Relationship between probability (x axis) odds and the log odds. Using log odds ensure a symmetric relationship with a zero midpoint (p = 0.5).

outcomes we have k-1 models determined by the cumulative probability in terms of $P(Y \le 1), \dots, P(Y \le k-1)$. We can manually calculate the odds ratio starting from the previous dataset.

```
# function to calculate odds
odds <- function(p) p / (1 - p)

# probability of Y = 1 for the group a
dat$yn <- as.integer(dat$y) # better as number here
(pia <- mean(dat$yn[dat$x == "a"] == 1))
#> [1] 0.38

# thus P(Y > 1) = 1 - pia
odds(pia)
#> [1] 0.6129032
# odds ratio Y = 1 a vs b
```

```
(p1b <- mean(dat$yn[dat$x == "b"] == 1))
#> [1] 0.14
odds(p1a) / odds(p1b)
#> [1] 3.764977
```

For example the probability of responding Y = 1 in the group "a" is 0.38 corresponding to an odds of 0.61. When calculating the odds ratio comparing "a" vs "b" for Y = 1 we obtain that the group "a" has 3.76 times the odds of responding Y = 1 compared to the group "b".

For an ordinal variable we have k levels and we can calculate k-1 cumulative probabilities (excluding the last associated with P=1). Thus we have k-1 odds ratio or what is called a cumulative odds ratio. Basically we repeat the previous steps but on the cumulative probability $P(Y \le k), k = 1, ..., k-1$. The three odds ratios are very similar correspond to the model estimate e_1^{β} , in R exp(fit\$beta) 4.92. In fact, the logit model essentially estimate the cumulative odds ratio.

```
# the dummy_ord() is a custom function creating k-1 dummy_ord() is a custom function creating k-1 dummy_ord()
# data.frame with the group (x) and the k-1 dummy variables
cum_p <- cbind(x = dat$x, dummy_ord(dat$y))</pre>
# calculating the cumulative probability for k-1 variables.
# Taking the average of a series of 0-1 is the same as computing the proportion of 1s.
(group_a \leftarrow apply(cum_p[cum_p$x == "a", -1], 2, mean))
#> y1vs234 y12vs34 y123vs4
     0.38
              0.60
                      0.86
(group_b \leftarrow apply(cum_p[cum_p$x == "b", -1], 2, mean))
#> y1vs234 y12vs34 y123vs4
    0.14 0.26 0.48
\# calculating k - 1 odds ratios on the cumulative probabilities as a/b
odds(group_a) / odds(group_b)
#> y1vs234 y12vs34 y123vs4
#> 3.764977 4.269231 6.654762
```

Proportional odds assumption

From the previous example, we noticed that the k-1 estimated odds ratios are different. However, increasing n will result in odds ratios that are very similar. This does not happen by chance but from an implicit (until now) assumption made by the CM called proportional odds (PO). Following again the taxonomy by Tutz (2022), each of the presented ordinal regression model has a basic version making the PO assumption. There are more advanced versions of the model relaxing this assumption completely (non proportional odds) and partially (partial proportional odds, Peterson & Harrell, 1990).

```
set.seed(2024)
# using same parameters as the previous simulation increasing n
n \leftarrow 1e5 \# very large n
x \leftarrow rep(c("a", "b"), each = n)
dat <- data.frame(x = x)</pre>
dat <- sim_ord_latent(~x, beta = b1, prob0 = probs, data = dat, link = "logit")</pre>
# same steps as before
cum_p <- cbind(x = dat$x, dummy_ord(dat$y))</pre>
group_a \leftarrow apply(cum_p[cum_p$x == "a", -1], 2, mean)
group_b <- apply(cum_p[cum_p$x == "b", -1], 2, mean)
# now odds are almost the same and equal to the simulated b1 = -1.09
odds(group_a) / odds(group_b)
#> y1vs234 y12vs34 y123vs4
#> 2.914579 2.985242 2.991333
log(odds(group_a) / odds(group_b))
#> y1vs234 y12vs34 y123vs4
#> 1.069725 1.093681 1.095719
```

Basically if we use the logit link function $g(\cdot)$, regression coefficients are interpreted as odds ratios. Given that we have k>2 alternatives we need k-1 equations. The PO assumption is formalized in Equation (4). The cumulative log odds ratio comparing $P(Y_k|x_0)$ with $P(Y_k|x_1)$ is the same regardless the specific threshold $(\beta_1=\beta_2\cdots=\beta_{k-1})$. The Figure (4) depicts the proportional odds assumption for the k-1 logistic curves both for probabilities and linear predictors η .

$$\operatorname{logit}(\frac{P(Y \leq 1|x_1)}{P(Y \leq 1|x_0)}) = \cdots = \operatorname{logit}(\frac{P(Y \leq k-1|x_1)}{P(Y \leq k-1|x_0)}) \tag{4}$$

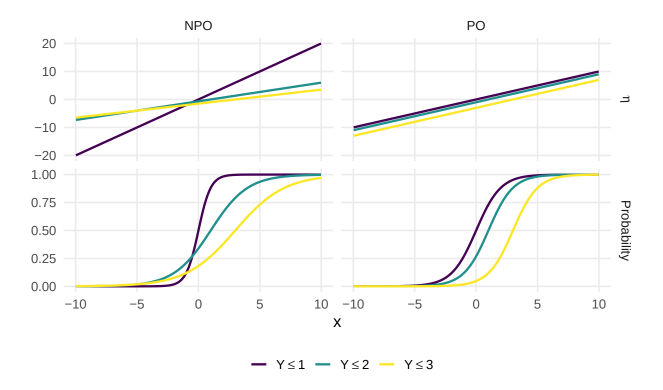


Figure 4. Example of proportional odds (PO, right) and non-proportional odds (NPO, left) for cumulative probabilities and the linear predictor η assuming a continuous predictor x. Data supporting the PO assumption shows an horizontal shift in the cumulative probability with the same slope while for NPO model slopes can be heterogeneous.

[Figure 4 about here]

The PO assumption is convenient because regardless of k, the β_j (j being the number of regression coefficients) effect is assumed to be the same. The model is more parsimonious compared to estimating k-1 coefficients for each β_j as in the multinomial regression or the non-proportional odds model.

At the same time, the PO assumption could be considered too strict. There are several methods for testing if data are supporting the PO assumption (see Liu, He, Tu, &

Tang, 2023 for an overview). Tutz and Berger (2020) suggested a trade-off between assuming/relaxing the PO assumption by fitting *location-shift* or *location-scale* models. Basically these methods should guarantee more flexibility in modelling the observed probabilities reducing the number of parameters. However, these methods and models are outside the scope of the tutorial. In the supplementary materials, we provide more details only about the *location-scale* model in terms of parameters interpretation and data simulation.

This can be shown by fitting the model with $\mathtt{clm}()$ that by default assume the PO and using the predicted probabilities to compute the odds ratios. In the following code we computed the odds ratio comparing x_a and x_b when $k \leq 1$ and $k \leq 2$. The odds ratios are exactly the same because we are using the predicted probabilities from the model (assuming PO).

```
# fitting the model
fit <- clm(y - x, data = dat, link = "logit")

# extracting the predicted probabilities for the two groups
pr <- predict(fit, data.frame(x = unique(dat$x)))$fit

# y <= 1
yia <- pr[1, 1]
yib <- pr[2, 1]

# y <= 2
y12a <- sum(pr[1, 1:2])
y12b <- sum(pr[2, 1:2])

# odds ratio y <= 1 (a vs b) VS y <= 2 (a vs b)
odds(y1a) / odds(y1b)

#> [1] 2.976954
odds(y12a) / odds(y12b)

#> [1] 2.976954
```

Probit Model and z scores

When assuming a standard Gaussian distribution we are fitting *probit* model. The main difference regards the parameters interpretation. In the logit model the β are the log odds ratio. For categorical variables they represents the increase in the log odds of moving from one level to the other while for numerical variables is the increase in the log odds for a unit increase in x.

For the *probit* model, the β is the increase in terms of z scores for a unit increase in x. This is very convenient especially for categorical variables because parameters can be interpreted as a Cohen's d like measure. Thinking about the latent distributions, the β is the shift in the latent mean comparing two or more groups or the slope of latent scores as a function of a numeric x. More formally the shift in the latent distribution is β/σ (for the *probit* model $\sigma=1$). The interpretation in terms of shifting the latent mean holds also for the logistic model. However, the standard deviation of the standard logistic regression is $\frac{\pi}{\sqrt{3}} \approx 1.81^3$. The β for the logistic distribution can be interpreted as the location shift of the latent logistic distribution by $\beta/(\frac{\pi}{\sqrt{3}})$ standard deviations (Agresti, 2010).

Proportional odds and probit model. The PO assumption is relevant only for the logit model. When using a different link function (e.g., probit) the assumption of a common β is called parallel slopes (see Equation (5)). Basically the difference in z scores for a unit increase in x (i.e., the slope) is the same regardless the threshold. As for the previous example, we can fit the probit model and manually calculate the difference in z scores.

$$z_{x_1-x_0} = \Phi(P \leq 1|x_1) - \Phi(P \leq 1|x_0) = \dots = \Phi(P \leq k-1|x_1) - \Phi(P \leq k-1|x_0) \eqno(5)$$

³ Actually the variance of the logistic distribution is $\frac{s^2\pi^2}{3}$ and the standard deviation $\frac{s\pi}{\sqrt{3}}$ where s is the scale of the distribution. For the standard logistic distribution s=1 (as for the standard normal distribution).

```
# fitting the model
fit <- clm(y - x, data = dat, link = "probit")

# extracting the predicted probabilities for the two groups
pr <- predict(fit, data.frame(x = unique(dat$x)))$fit

# y <= 1
y1a <- pr[1, 1]
y1b <- pr[2, 1]

# y <= 2
y12a <- sum(pr[1, 1:2])
y12b <- sum(pr[2, 1:2])

# z score difference y <= 1 (a vs b) VS y <= 2 (a vs b)
qnorm(y1a) - qnorm(y1b)

#> [1] 0.6523531
qnorm(y12a) - qnorm(y12b)
#> [1] 0.6523531
```

Simulating data

In this tutorial we present two method for simulating ordinal data. Simulating data is a powerful strategy to understand the model (DeBruine & Barr, 2021) and estimate statistical properties (e.g., power or type-1 error). The first simulation method calculate the probabilities of each Y level as a function of predictors and generate data from a multinomial distribution. The second method simulate data using the the latent formulation. Whenever random number generation occurs, it is appropriate to set a seed (set.seed() function, in this case we use set.seed(2024)). Running again the same code will produce the same result. A general simulation approach concerns generating a dataset from the assumed data generation process, fitting the statistical model and assessing the recovery of simulated parameters. To check the simulation approach, using a large sample size produce estimations with small sampling error. A limited sample size even when fitting the true model will produce variable estimations.

Simulating from a multinomial distribution. For the first method we need to calculate $g^{-1}(\eta)$ as a function of predictors and then sample from a multinomial (more specifically categorical) distribution using the sample() function in R. This method is similar to the general way of simulating data for a generalized linear model (see the supplementary materials for a general overview).

As a simple example, let's simulate two groups with n = 100 participants responding to an item (Y) with k = 4 ordered options. We simulate the second group having higher probability of responding higher categories of Y.

As explained in the previous sections, we can summarise the effect size of a CM (assuming PO) using a single $\beta = \log(OR)$. We can assume that the odds ratio is OR = 2. For simplicity, the probabilities of the first group x = 0 are uniform thus $P(Y = 1|x_0) = \dots P(Y = k|x_0) = 1/k$. The following code summarise the first steps of the simulation. Basically we define the simulation parameters, we calculate the k-1 thresholds α and we apply Equations (1) and (2). In this way we calculated the true probability of each Y level for the two groups.

An important step is converting from α to probabilities or the opposite. This steps are implemented in the alpha_to_prob() and prob_to_alpha() functions that given the input return thresholds or probabilities.

```
k <- 5

(p <- rep(1/k, k)) # uniform

#> [1] 0.2 0.2 0.2 0.2 0.2 0.2

names(p) <- paste0("y", 1:k)

(alpha <- prob_to_alpha(p, link = "logit")) # or prob_to_alpha(p, "probit")

#> y1  y2  y3  y4

#> -1.3862944 -0.4054651 0.4054651 1.3862944

alpha_to_prob(alpha, link = "logit")

#> [1] 0.2 0.2 0.2 0.2 0.2 0.2
```

```
## SIMULATION PARAMETERS
N <- 100 # sample size
or <- 2 # odds ratio
k \leftarrow 4 # number of ordinal alternatives
prob0 <- rep(1/k, k) # probabilities for the first group</pre>
alpha <- prob_to_alpha(prob0, link = "logit")</pre>
dat \leftarrow data.frame(x = rep(c(0, 1), each = N/2))
## LINEAR PREDICTOR
\# calculate linear predictor using equation 1 obtaining k - 1 equations
lp <- lapply(alpha, function(a) a - log(or) * dat$x)</pre>
names(lp) <- sprintf("lp_leq%s", 1:(k - 1)) # giving appropriate names</pre>
lp <- data.frame(lp)</pre>
head(lp)
#> lp_leq1 lp_leq2 lp_leq3
#> 1 -1.098612
                 0 1.098612
#> 2 -1.098612
                  0 1.098612
#> 3 -1.098612
                  0 1.098612
#> 4 -1.098612
                 0 1.098612
#> 5 -1.098612
                  0 1.098612
#> 6 -1.098612
                 0 1.098612
## CUMULATIVE PROBABILITIES
# apply the inverse of the link function (invlogit) to calculate cumulative probabilities
cump <- lapply(lp, plogis)</pre>
cump <- data.frame(cump)</pre>
names(cump) <- sprintf("cump_leq%s", 1:(k - 1)) # giving appropriate names</pre>
head(cump)
#> cump_leq1 cump_leq2 cump_leq3
#> 1
        0.25
                  0.5 0.75
                  0.5 0.75
#> 2
        0.25
        0.25
#> 3
                  0.5
                          0.75
        0.25
                  0.5
                            0.75
#> 4
#> 5
        0.25
                  0.5
                            0.75
#> 6
         0.25
                  0.5
                            0.75
## PROBABILITIES OF Y
```

```
# for each row, we can calculate P(Y = k) using equation 2
\# P(Y = 1) = P(Y \le 1)
\# P(Y = 2) = P(Y \le 2) - P(Y \le 1)
\# P(Y = 3) = P(Y \le 3) - P(Y \le 2)
\# P(Y = 4) = 1 - P(Y \le 3)
\# adding a columns of 0 and 1, then diff() for adjacent differences
cump <- cbind(0, cump, 1)</pre>
p <- apply(cump, 1, diff, simplify = FALSE)</pre>
p <- data.frame(do.call(rbind, p)) # collapse list of rows into a dataframe
names(p) <- sprintf("p%s", 1:k) # giving appropriate names</pre>
head(p)
#> p1 p2 p3 p4
#> 1 0.25 0.25 0.25 0.25
#> 2 0.25 0.25 0.25 0.25
#> 3 0.25 0.25 0.25 0.25
#> 4 0.25 0.25 0.25 0.25
#> 5 0.25 0.25 0.25 0.25
#> 6 0.25 0.25 0.25 0.25
# probabilities for id = 1 (x = 0) and id = 51 (x = 1)
p[c(1, 51),]
#> p1
                    p2 p3 p4
#> 1 0.2500000 0.2500000 0.2500000 0.25
#> 51 0.1428571 0.1904762 0.2666667 0.40
## CUMULATIVE ODDS RATIO
# calculate the (cumulative) odds ratio
x0 <- cump[1, 2:k]</pre>
x1 \leftarrow cump[51, 2:k]
\# this is the same as the or (the b1) and we are assuming POA
odds(x0) / odds(x1)
#> cump_leq1 cump_leq2 cump_leq3
#> 1 2 2
```

```
## SAMPLING FROM THE MULTINOMIAL DISTRIBUTION
set.seed(2024)
\hbox{\it\# example of a random Y outcome based on the probabilities}
sample(x = 1:k, size = 1, prob = p[1, ])
#> [1] 1
# we can apply it to the full dataset, this step lead to different results
\hbox{\it\# each time we run the code because we are sampling from a distribution}
dat$y <- apply(p, 1, function(ps) sample(1:k, size = 1, prob = ps))</pre>
head(dat)
#> x y
#> 1 0 3
#> 2 0 4
#> 3 0 4
#> 4 0 3
#> 5 0 4
#> 6 0 3
# let's compute the observed probabilities, to be compared to the true
# probabilities
# observed
(op <- prop.table(table(dat$x, dat$y), margin = 1))</pre>
        1 2 3 4
#> 0 0.26 0.22 0.26 0.26
#> 1 0.10 0.18 0.30 0.42
# true (just selecting a row from x = 0 and x = 1)
p[c(1, 51),]
          p1
                    p2 p3 p4
#> 1 0.2500000 0.2500000 0.2500000 0.25
#> 51 0.1428571 0.1904762 0.2666667 0.40
\# similarly we can compute the observed cumulative odds ratios
(cum_op <- apply(op, 1, cumsum))</pre>
```

```
dat$y <- ordered(dat$y) # make an ordered factor in R where 1 < 2 < 3 < 4
fit <- clm(y ~ x, data = dat, link = "probit")</pre>
summary(fit)
\# formula: y \sim x
#> data: dat
#>
#> probit flexible 100 -132.59 273.17 5(0) 2.86e-09 1.7e+01
#>
#> Coefficients:
\#> Estimate Std. Error z value Pr(>|z|)
#> x 0.5183 0.2197 2.359 0.0183 *
#> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
#>
#> Threshold coefficients:
#> Estimate Std. Error z value
#> 3|4 0.68706 0.17462 3.935
```

We can generalize the previous workflow as:

- 1. Define simulation parameters, baseline probabilities (prob0), sample size, etc.
- 2. Define regression coefficients β and calculate the k-1 linear predictors (η) using the equations

- 3. Apply the inverse of the link function $g^{-1}(\eta)$ on the linear predictor and calculate the cumulative probabilities $p(y \le 1|x), p(y \le 2|x), ..., p(y \le k-1|x)$.
- 4. Calculate the probabilities of k outcomes
- 5. Sample n outcomes from a multinomial distribution, choosing between k alternatives using the calculated probabilities
- 6. Fit the appropriate model using ordinal::clm()

Simulating from the latent distribution. A more efficient way to simulate an ordinal outcome is using the latent formulation of the model. This require simulating a standard linear regression using the appropriate data generation function (logistic or normal) and the cutting the latent values according to the thresholds α . The workflow is slightly different compared to the previous approach.

- 1. Define simulation parameters as in the previous simulation. prob0 are the probabilities when all predictors X are zero.
- 2. Define regression coefficients β and calculate the linear predictor η using the Equation (3)
- 3. Add the random errors ϵ_i sampling from the chosen distribution (logistic or normal)
- 4. Cut the latent variable into k areas using the thresholds α and assign the corresponding ordinal value. This can be done using the cut() or the findInterval() functions.

The Figure 5 depicts the simulated Y^* and the corresponding ordinal value. As for the previous simulation we can fit the model using clm() and check the estimated parameters.

```
## SIMULATION PARAMETERS

set.seed(2024)
N <- 1e3 # sample size</pre>
```

```
or <- 4 # odds ratio, higher here just for a more clear plot
k \leftarrow 4 # number of ordinal alternatives
probs0 <- rep(1/k, k) # probabilities for the first group</pre>
alpha <- prob_to_alpha(probs0, link = "logit")</pre>
dat \leftarrow data.frame(x = rep(c(0, 1), each = N/2))
## LINEAR PREDICTOR
# calculate the linear predictor using the model equation
dat$lp <- log(or) * dat$x</pre>
# add the random part by sampling errors from a standard logistic (or normal) distribution
dat$ystar <- dat$lp + rlogis(N, location = 0, scale = 1)</pre>
# cut the latent distribution. The + 1 because the first category is 0 by default.
dat$y <- findInterval(dat$ystar, alpha) + 1</pre>
head_tail(dat, n = 3)
      x lp
                   ystar y
     0 0.000000 1.6356528 4
#> 1
#> 1000 1 1.386294 2.0325030 4
```

[Figure 5 about here]

```
dat$y <- ordered(dat$y) # make an ordered factor in R where 1 < 2 < 3 < 4
fit <- clm(y ~ x, data = dat, link = "logit")
summary(fit)

#> formula: y ~ x

#> data: dat

#>
#> link threshold nobs logLik AIC niter max.grad cond.H

#> logit flexible 1000 -1226.66 2461.32 4(0) 2.25e-07 1.7e+01

#>
#> Coefficients:
#> Estimate Std. Error z value Pr(>|z|)
#> x 1.4078 0.1233 11.41 <2e-16 ***</pre>
```

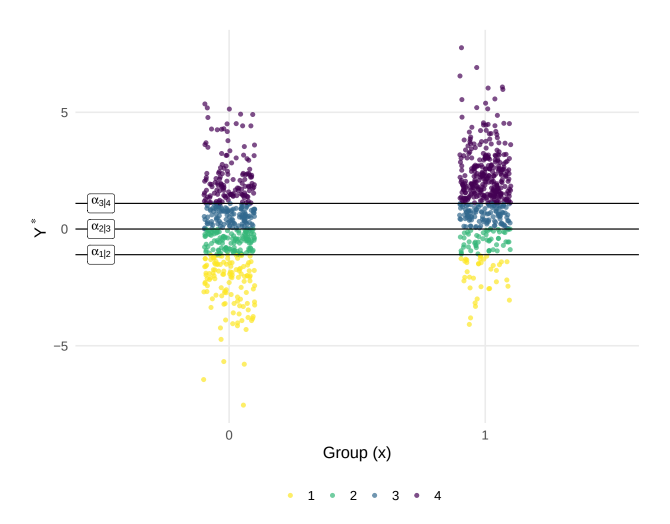


Figure 5. Example of cutting a latent variable for k = 4 ordinal outcomes with the effect of a binary predictor x. The thresholds are fixed and the difference between the latent means of the two groups increase the number of points (thus probability) in higher outcomes.

The simulation using the latent formulation of the model is implemented in the $sim_ord_latent()$ function. Basically, we define the dataset dat with predictors. Then the model formula is specified within the function as $location = \sim along$ with the vector of regression coefficients (beta), baseline probabilities (prob0) and the link function. The function return a dataset with the simulated Y and the latent variable (this is possible only because we simulate the data, otherwise Y^* is not observed by definition).

```
set.seed(2024)
N <- 100 \# sample size
or <- 4 \# odds ratio, higher here just for a more clear plot
k \leftarrow 4 # number of ordinal alternatives
probs0 <- rep(1/k, k) # probabilities for the first group
alpha <- prob_to_alpha(probs0, link = "logit")</pre>
dat \leftarrow data.frame(x = rep(c(0, 1), each = N/2))
# same as the previous simulation
dat <- sim_ord_latent(location = ~x, beta = log(or), prob0 = probs0, data = dat, link = "logit")</pre>
head_tail(dat, n = 3)
      x y
#> 1 0 4 1.6356528
#> 2  0  2  -0.7497878
#> 3 0 3 0.7554420
#> 100 1 4 1.6703959
```

Choosing parameters values.

Thresholds α . The previous simulation can be easily extended by adding more predictors and their interactions. The crucial part is setting appropriate and empirically meaningful parameters. Thresholds α are usually not of main interest (but see the location-shift models Tutz, 2022) and can be considered as intercepts in standard linear regression. The thresholds are quantiles of the latent distribution that produced certain k probabilities. To set meaningful α values we can convert probabilities into thresholds (alpha_to_prob()). The function show_alpha() produce a meaningful visual representation of using a specific set of thresholds (see Figure 6). In regression terms, the thresholds determine the probabilities when x = 0 (as the intercept). Thus with two groups for example, the thresholds are the k probabilities of the ordinal variable Y for the reference group.

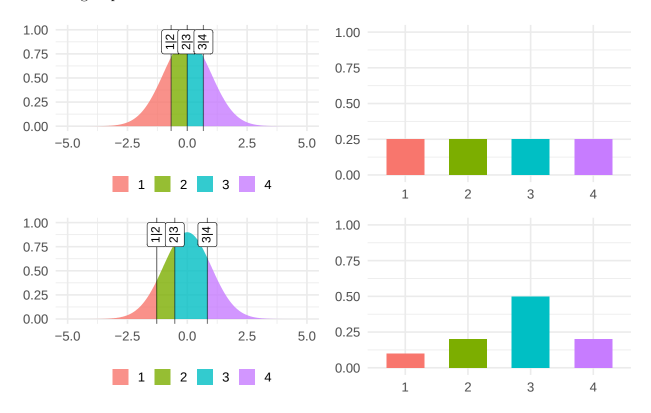


Figure 6. Example of the $show_alpha()$ function. On the left the assumed latent variable and k-1 thresholds with the corresponding probabilities on the right.

Regression coefficients. For probit models we can set the β_j to be in standardized (i.e., Cohen's d-like) units. For a categorical variable as the previous example with the group, β_j is the degree of separation in standard deviation unit between the two latent distributions. For logit models we can set the odds ratio. Meaningful odds ratios can be derived from previous literature, meta-analyses or converting to other effect sizes. For example, Sánchez-Meca, Marín-Martínez, and Chacón-Moscoso (2003) proposed some equations to convert between odds ratios and Cohen's d. Using their approach, a Cohen's d = 0.5 usually considered a plausible medium effect size corresponds to and odds ratio of ≈ 2.47 .

We can also calculate and plot the predicted probabilities (i.e., $g^{-1}(\eta)$) given the predictors and the chosen regression coefficients. In this way we can try different values and see if predicted probabilities are plausible or not. The cat_latent_plot() and num_latent_plot() functions can be for respectively a categorical (Figures 7) and numerical predictor (Figures 8).

[Figure 7 about here]

[Figure 8 about here]

2x2 interaction

A common research design could be a 2x2 factorial design. In this example we have two main effects and the interaction. By default R use dummy coding but setting sum-to-zero contrasts (e.g., 0.5 and -0.5) for a factorial design is convenient. In this way β_1 will be the main effect of X_1 , β_2 the main effect of X_2 and β_3 the interaction (thus the difference of differences). Equation (6) depict the model formula.

$$P(Y_i \le k) = g^{-1}[\alpha_k - (\beta_1 X_{1_i} + \beta_2 X_{2_i} + \beta_3 X_{1_i} X_{2_i})] \tag{6}$$

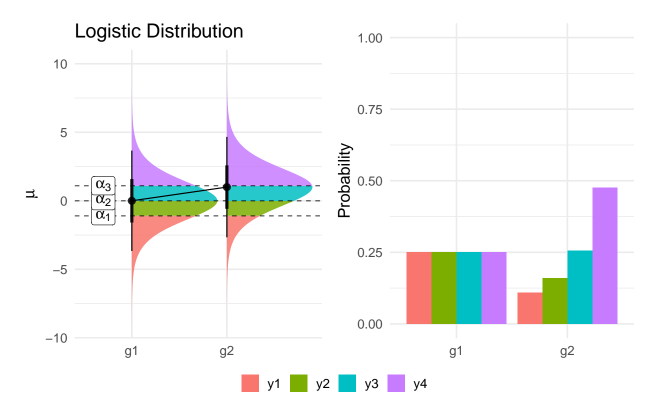


Figure 7. Example of the cat_latent_plot() function depicting the effect of a categorical predictor. On the left the shift in the latent mean and on the right the impact on the expected probabilities. The plot can be created using cat_latent_plot(m = c(0, 0.5), s = 1, probs = rep(1/4, 4), link = "logit", plot = "both").

```
set.seed(2024)
n <- 100
k <- 3
betas <- c(b1 = 0, b2 = 1, b3 = 0.5) # b1 = main effect X1, b2 = main effect X2, b3 = interaction

dat <- expand.grid(x1 = c("a", "b"), x2 = c("c", "d"), n = 1:n)
dat$x1 <- factor(dat$x1)
dat$x2 <- factor(dat$x2)

# sum to 0 coding
contrasts(dat$x1) <- c(0.5, -0.5)
contrasts(dat$x2) <- c(0.5, -0.5)
probs0 <- rep(1/k, k)</pre>
```

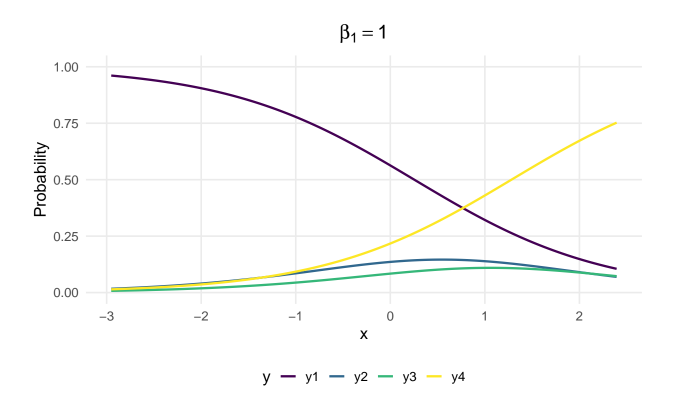


Figure 8. Example of the num_latent_plot() function depicting the effect of a continuous predictor on the expected probabilities. β_1 is the regression coefficient. The plot can be created using num_latent_plot(x = runif(100), b1 = 1, probs = c(0.6, 0.2, 0.1, 0.1), link = "probit")

```
dat <- sim_ord_latent(~ x1 * x2, beta = betas, prob0 = probs0, link = "probit", data = dat)
fit <- clm(y ~ x1 * x2, data = dat, link = "probit")</pre>
summary(fit)
\# formula: y \sim x1 * x2
           dat
#> data:
#>
#> link threshold nobs logLik AIC niter max.grad cond.H
#> probit flexible 400 -390.29 790.57 4(0) 2.20e-07 2.1e+01
#>
#> Coefficients:
          Estimate Std. Error z value Pr(>|z|)
#> x11
           -0.02969
                      0.11625 -0.255
                                         0.798
           1.15703
                      0.11935 9.695
                                        <2e-16 ***
#> x11:x21 0.29058
                      0.23254
                               1.250
                                         0.211
```

```
#> ---
#> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
#>
#> Threshold coefficients:
#> Estimate Std. Error z value
#> 1/2 -0.50503     0.06932  -7.286
#> 2/3     0.46102     0.06900     6.681
```

The parameters interpretation is the same as introduced in the general case. The thresholds α are fixed representing the probabilities P(Y=k) when all predictors are zero. In this case by doing alpha_to_prob(fit\$alpha, link = "probit") we should recover the probs0 vector. x1 is the main effect thus the difference in z scores between a and b averaging over x2. The same holds for x2. x1:x2 is the interaction thus difference of differences in z scores (see Figure 9).

[Figure 9 about here]

Numerical by categorical interaction

Another common scenario is the interaction between a numerical variable x and a categorical variable g. For simplicity we simulate the factor with two levels and the numerical variable sampled from an uniform distribution between 0 and 1.

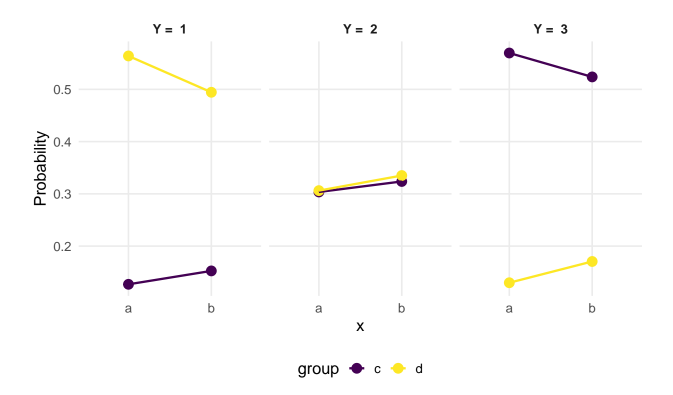


Figure 9. Results from the simulated model with a 2x2 interaction between categorical predictors. The plot shows the predicted probabilities for each ordinal outcome as a function of the two predictors. A similar plot can be produced using plot(ggeffects::ggpredict(fit, terms = c("x1", "x2"))).

```
#> data: dat

#>

#> link threshold nobs logLik AIC niter max.grad cond.H

#> probit flexible 100 -101.63 213.26 5(0) 3.01e-08 1.0e+02

#>

#> Coefficients:

#> Estimate Std. Error z value Pr(>|z|)

#> x 0.68616 0.40959 1.675 0.0939 .

#> g1 0.07207 0.45768 0.157 0.8749

#> x:g1 1.21471 0.81881 1.484 0.1379

#> ---

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

#>

#> Threshold coefficients:
```

Again, x is the slope of the numerical predictor averaging over g (given that g has been coded with sum-to-zero contrasts) thus the increase in z scores for a unit increase in x. g1 is the main effect of the factor evaluated when x=0 (centering x will change the parameter interpretation). The x:g1 is the slopes difference the two groups. Figure 10 depicts the model results.

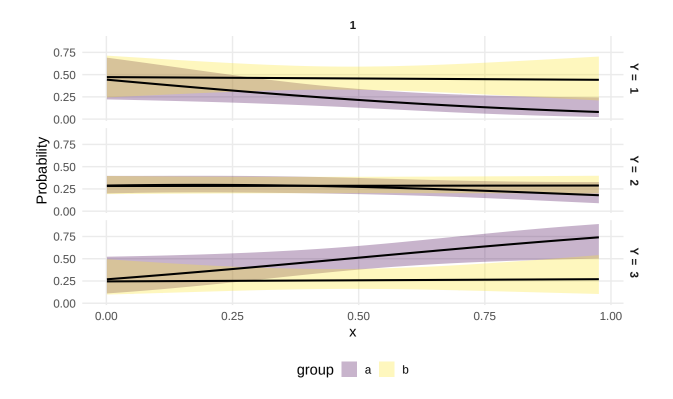


Figure 10. Results from the simulated model with the interaction between a continuous and categorical variables. The plot shows the predicted probabilities with 95% confidence intervals as a function of predictors. A similar plot can be produced using plot(ggeffects::ggpredict(fit, terms = c("x[all]", "g")))

Power Analysis

In this section we introduce how to estimate the statistical power. Power analysis is a crucial and often necessary step when planning a confirmatory study (Lakens, 2022) or preparing a pre-registration or registered report (Chambers & Tzavella, 2022). Furthermore, statistical power for complex models can only be estimated using Monte Carlo simulations.

Formally the power is defined as the probability of correctly rejecting the null hypothesis H_0 . For simple cases (e.g., a t-test) the power can be calculated analytically while in complex cases simulating data is the best approach. We present very general example that, as for the previous simulations, can be easily extended. For example, by simulating a model fixing a specific parameter to zero, we can estimate the type-1 error rate (wrongly rejecting H_0) or assessing an eventual bias in model estimates.

The general workflow for a power simulation can be summarized as:

- 1. Specify the research design, e.g., 2x2 factorial design
- 2. Specify the effect/parameter of interest, e.g., the interaction effect
- 3. Define the simulation conditions, e.g., a range of sample size, effect size, etc.
- 4. Implement one of the simulation workflow described in the previous sections
- 5. Repeat the simulation a large number of times (e.g, 10000) and store the relevant values from each simulation
- 6. Summarise the simulation results

We can estimate the power of detecting a group difference of d = 0.4 (assuming a probit model). Participants respond to an ordinal variable with k = 5. The simulation is performed using a for loop that repeat 1000 times the data simulation, model fitting and extract the p-value for the β_1 . The power is then estimated as the number of p-values lower than the critical level over the number of simulations.

```
set.seed(2024)
n <- 40 # sample size
k <- 5 # number of ordinal variables
d <- 0.4 # effect size (i.e., our regression coefficients)
nsim <- 1e3 # higher is better, here using 1000 for an example
probs0 \leftarrow rep(1/k, k)
alpha <- 0.05 # critical alpha
p <- rep(NA, nsim) # preallocation to improve loop computational efficiency
dat <- data.frame(group = rep(c("a", "b"), each = n)) # data frame</pre>
head_tail(dat, n = 3)
      group
#> 1
#> 2
#> 3
#> 78
#> 79
#> 80
for(i in 1:nsim){
  sim <- sim_ord_latent(~group, beta = d, prob0 = probs0, link = "probit", data = dat)</pre>
 fit <- clm(y ~ group, data = sim, link = "probit")</pre>
  p[i] <- summary(fit)$coefficients["groupb", "Pr(>|z|)"] # extract the pvalue
}
# estimate the power
mean(p <= alpha, na.rm = TRUE)</pre>
#> [1] 0.418
```

Despite useful, calculating power curves instead a single value is more informative. For this reason we repeat the previous simulation but for different sample sizes. Figure 11 depict the power curve resulting from the simulation.

```
set.seed(2024)

n <- c(20, 40, 60, 100, 200)
power <- rep(NA, length(n))

for(i in 1:length(n)){</pre>
```

```
p <- rep(NA, nsim) # preallocation for speed

dat <- data.frame(group = rep(c("a", "b"), each = n[i]))

for(j in 1:nsim){
    sim <- sim_ord_latent(~group, beta = d, prob0 = probs0, link = "probit", data = dat)
    fit <- clm(y ~ group, data = sim, link = "probit")
    p[j] <- summary(fit)$coefficients["groupb", "Pr(>|z|)"]
}

power[i] <- mean(p <= alpha)
}

power

#> [1] 0.225 0.412 0.522 0.750 0.964
```

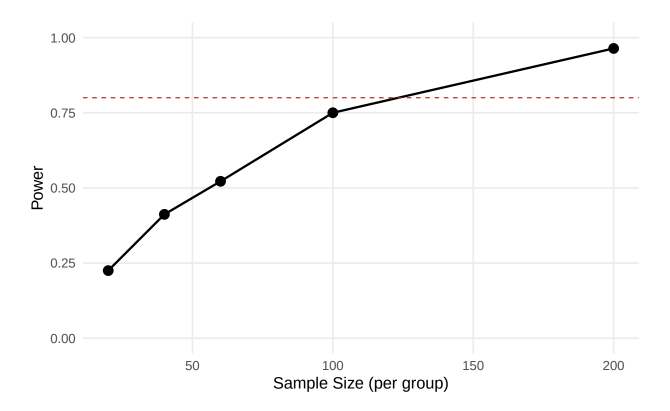


Figure 11. Results from the power analysis. The x-axis represents the sample size (per group) and the y axis the estimated power. The red dotted line highlight the 80% power level.

[Figure 11 about here]

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