

# ordPens: An R package for Selection, Smoothing and Principal Components Analysis for Ordinal Variables

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## Summary

Ordinal data are a common case in applied statistics. In order to incorporate the ordinal scale level, among other things, regularization techniques are often suggested in the literature. In particular, penalization approaches for smoothing and selection when dealing with Likert-type data – which are by no means restricted to Likert scale – are commonly proposed. **ordPens** is a package in the R programming language (R Core Team, 2021) and provides several penalty approaches for ordinal predictors in regression models and ordinal variables for principal component analysis (PCA).

The main idea behind smoothing in the regression context is to maximize the penalized log-likelihood by introducing a penalty term and a tuning parameter controlling the amount of penalty. Different types of penalization can be considered, depending on whether to achieve smoothing, selection or clustering of variables. Smoothing only can be done by penalizing the sum of squared differences of adjacent coefficients. A modified group lasso based on a difference penalty can be used for selection. Clustering/fusion of categories can be achieved by the fused lasso penalizing absolute differences by using the  $L_1$ -norm.

## Statement of Need

As suggested by Tutz & Gertheiss (2014) and Tutz & Gertheiss (2016), selection, and/or smoothing/fusing of ordinally scaled independent variables shall be done using a modified group lasso or generalized ridge penalty when dealing with ordinally scaled predictors in regression analysis. The penalized log-likelihood to be maximized takes the form  $l_p(\beta) = l(\beta) - \lambda J(\beta)$ , with  $\beta$  corresponding to the vector of regression parameters,  $\lambda$  representing the smoothing parameter and  $J(\cdot)$  being the penalty function.

**ordPens** (Gertheiss & Hoshiyar, 2021) offers various tools for data analysis of ordinally scaled data. The package attacks the afore mentioned tasks and offers penalized regression for smoothing, selection and fusion. Specifically, the function **ordSmooth()** for smoothing only incorporates the generalized ridge penalty

$$J(\beta) = \sum_{s=1}^p \beta_s^T D_{d,s}^T D_{d,s} \beta_s,$$

with  $D_{d,s}$  being the matrix generating differences of order  $d$  and  $\beta_s^T = (\beta_{s1}, \dots, \beta_{sk_s})$  being the parameter vector linked to the  $s$ th (dummy-coded) predictor with categories  $1, \dots, k_s$ . The **ordSelect()** function performs smoothing and selection by adopting a modified group lasso penalty based on differences of the form

$$J(\beta) = \sum_{s=1}^p \sqrt{k_s} \sqrt{\beta_s^T D_{d,s}^T D_{d,s} \beta_s}.$$

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Clustering of categories is done by the function `ordFusion()`, which uses a fused lasso penalty based on differences of first order:

$$J(\beta) = \sum_{s=1}^p \sum_{j=2}^{k_s} |\beta_{sj} - \beta_{s,j-1}|.$$

For more information on the original group lasso (for nominal predictors and grouped variables in general), see Meier, Van De Geer, & Bühlmann (2008) and Yuan & Lin (2006). For details on the fused lasso, see Tibshirani, Saunders, Rosset, Zhu, & Knight (2005). In the case of smoothing only, the package includes auxiliary functions such that `mgcv::gam()` (Wood, 2008, 2017) can be used for fitting generalized linear and additive models with first- and second-order ordinal smoothing penalty as well as built-in smoothing parameter selection. Also, `mgcv` tools for further statistical inference can be used, see Gertheiss, Scheipl, Lauer, & Ehrhardt (2021) for details. Furthermore, testing for differences in the means, known as analysis of variance (ANOVA) is provided for ordered factors by the function `ordAOV()` penalizing (squared) differences of adjacent means. Testing for differentially expressed genes, when analyzing microarrays of gene expression data, is incorporated by the function `ordGene()`. Technical details can be viewed from Gertheiss (2014) and Sweeney, Crainiceanu, & Gertheiss (2016), respectively.

If, in contrast, dimension reduction is desired in an unsupervised way, principal components analysis can be applied to ordinal data as well. However, those data are usually either treated as numeric implying linear relationships between the variables at hand, or non-linear PCA is applied where the obtained coefficients are sometimes hard to interpret. Note that in IBM SPSS Statistics (Version 25.0), for instance, there is an option available for smoothing quantifications by the use of spline functions, which, however, limits the type of functions that can be fitted when using a small number of knots and a suitable choice may be challenging for the (inexperienced) user. On the other hand, as splines are defined on interval scale whereas ordinal variables can only take some discrete values, the usage of spline functions may be seen as unnecessarily complex for scaling ordinal data. To incorporate the ordinal scale level, the concept of penalization can also be adapted here, as suggested in Hoshiyar, Kiers, & Gertheiss (2021). Penalized non-linear principal components analysis for ordinal variables is incorporated in the function `ordPCA()` using a second-order difference penalty. In addition, the function provides performance evaluation and selection of an optimal penalty parameter using k-fold cross-validation. Also, the option of both non-monotone effects and incorporating constraints enforcing monotonicity is provided. Penalized non-linear PCA therefore serves as an intermediate between the standard methods typically used so far (see above). The new approach offers both better interpretability as well as better performance on validation data.

## Availability

The R package `ordPens` is publicly available on CRAN and Github, where issues can be opened. `ordPens` is licensed under the GPL-2 General Public License. Documentation and examples are contained in the package manual, which can be found on CRAN.

To install `ordPens`, simply run:

```
install.packages("ordPens")
```

For penalized regression and ordinal ANOVA see also `vignette("ordPens", package = "ordPens")`. Penalized non-linear PCA is also documented in detail and can be accessed via `vignette("ordPCA", package = "ordPens")`.

## ordPens in action

This example illustrates penalized non-linear PCA on the so-called brief ICF core set on Chronic Widespread Pain (CWP) consisting of 26 ordinal scaled variables. Details on the data can be found in Gertheiss, Hogger, Oberhauser, & Tutz (2011); type also `?ICFCoreSetCWP`. Analysis of the so-called comprehensive core set for CWP, consisting of 67 ICF variables, is found in Hoshiyar et al. (2021). Note that the `ordPCA` vignette also analyzes the comprehensive core set.

```

library(ordPens)

# load ICF data & code adequately
data(ICFCoreSetCWP)
H <- ICFCoreSetCWP[, 1:67] + matrix(c(rep(1,50), rep(5,16), 1),
                                   nrow(ICFCoreSetCWP), 67, byrow = TRUE)

# select brief core set variables
brief <- c("b130","b134","b140","b147","b152","b1602","b280","b455","b730","b760",
          "d175","d230","d240","d430","d450","d640","d760","d770","d850","d920",
          "e1101","e310","e355","e410","e420","e570")
H <- H[, brief]
xnames <- names(H)

# ordinal penalized PCA
icf_pca1 <- ordPCA(H, p = 2, lambda = c(5, 0.5, 0.001), maxit = 100,
                  crit = 1e-7, qstart = NULL, Ks = c(rep(5,20), rep(9,6)),
                  constr = c(rep(TRUE,20), rep(FALSE,6)), CV = FALSE, k = 5)

# 5-fold cross-validation
lambda <- 10^seq(4, -4, by = -0.1)
set.seed(1234)
cvResult <- ordPCA(H, p = 2, lambda = lambda, Ks = c(rep(5,20), rep(9,6)),
                  constr = c(rep(TRUE,20), rep(FALSE,6)),
                  CV = TRUE, k = 5, CVfit = FALSE)

# plotting results for selected variables
par(mfrow = c(2,3))
for(i in which(xnames=="b280" | xnames=="d450" | xnames=="e1101" | xnames=="e410")){
  plot(icf_pca1$qs[[i]][,3], type = "b", xlab = "category", ylab = "quantification",
       col = 1, ylim = range(icf_pca1$qs[[i]]), main = xnames[i], bty = "n", xaxt = "n")
  lines(icf_pca1$qs[[i]][,2], type = "b", col = 2, lty = 2, pch = 2, lwd = 2)
  lines(icf_pca1$qs[[i]][,1], type = "b", col = 3, lty = 3, pch = 3, lwd = 2)
  axis(1, at = 1:length(icf_pca1$qs[[i]][,1]))
}

plot(log10(lambda), apply(cvResult$VAFtrain, 2, mean), type = "l",
     xlab = expression(log[10](lambda)), ylab = "VAF", main = "training data",
     cex.axis = 1.2, cex.lab = 1.2)

plot(log10(lambda), apply(cvResult$VAFtest, 2, mean), type = "l",
     xlab = expression(log[10](lambda)), ylab = "VAF", main = "validation data",
     cex.axis = 1.2, cex.lab = 1.2)
abline(v = log10(lambda)[which.max(apply(cvResult$VAFtest, 2, mean))], lty = 2)

```

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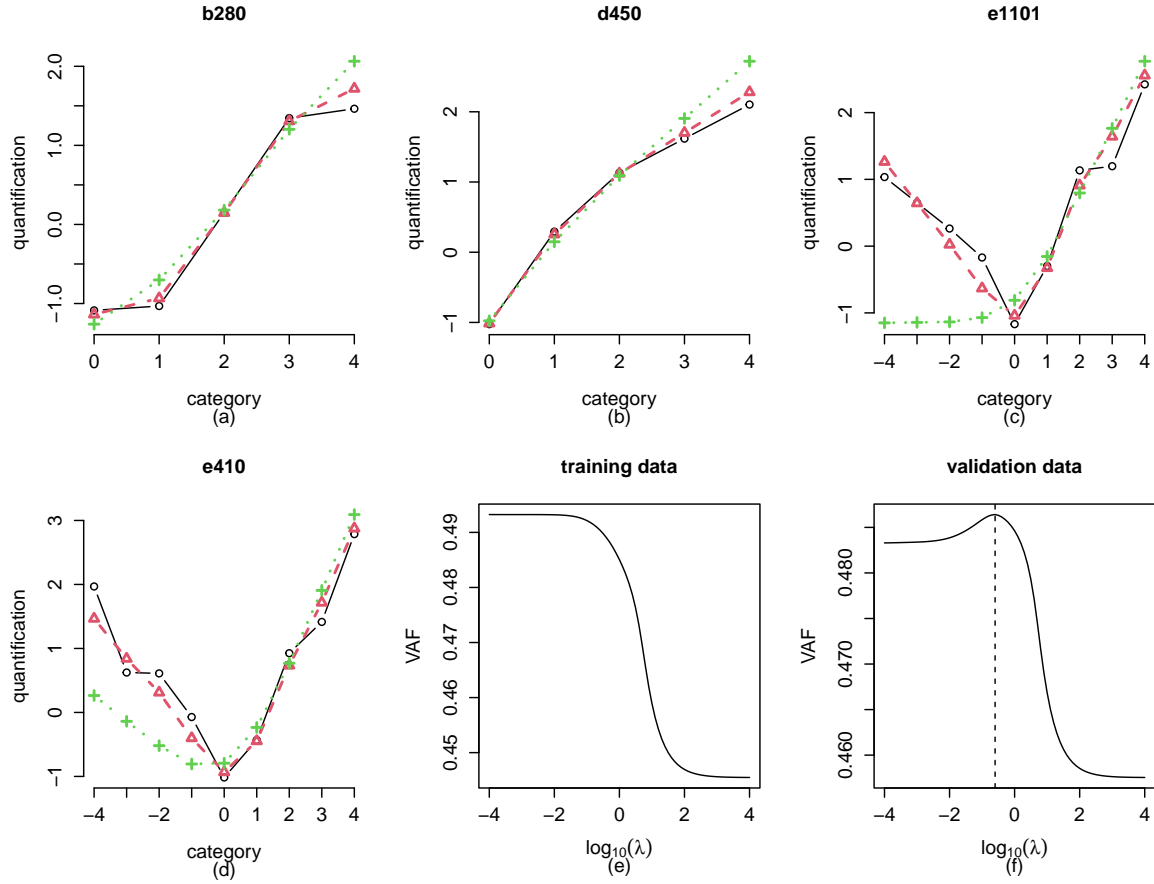


Figure 1: Category quantifications/scores for  $\lambda \rightarrow 0$  (solid black),  $\lambda = 0.5$  (dashed red),  $\lambda = 5$  (dotted green) (a)–(d); VAF by the first 5 principal components: (e) training data, (f) validation data with optimal  $\lambda$  (dashed line).

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