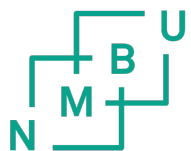


RAJU RIMAL*, TRYGVE ALMØY & SOLVE SÆBØ

EXPLORATION OF MULTI-RESPONSE
MULTIVARIATE METHODS



AN APPLICATION OF SIMREL
BIAS @ KBM, NMBU
<https://www.mathatistics.com/>
2019, July

Raju Rimal*, Trygve Almøy & Solve Sæbø: *Exploration of Multi-Response Multivariate Methods*, An application of simrel, © 2019, July. WEBSITE:

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The titlepage reproduces an engraving of Maurits Cornelis Escher, titled *Plane Filling with Birds* (the picture is obtained from <http://www.mcescher.com/>).

ABSTRACT

This section belongs to abstract. Write awesome abstract.

SUMMARY

This is summary section of the thesis.

ACKNOWLEDGEMENTS

I wish first of all to thank the members of the Staff of the Italian TeX{} and L^AT_EX{} User Group, in particular Prof.~Enrico Gregorio, for their invaluable aid during the writing of this work, the detailed explanations, the patience and the precision in the suggestions, the supplied solutions, the competence and the kindness. Thanks also to all the people who have discussed with me on the forum of the Group, prodigal of precious observations and good advices.

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1

INTRODUCTION

Citing Rimal et al. [2] in this text.

2 | METHODS

This is the introduction section. It contains background subsection

2.1 BACKGROUND

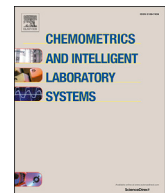
Here I will reference to [?] I hope this will render properly.

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- [1] Raju Rimal, Trygve Almøy, and Solve Sæbø. A tool for simulating multi-response linear model data. *Chemometrics and Intelligent Laboratory Systems*, 176:1 – 10, 2018. ISSN 0169-7439. doi: <https://doi.org/10.1016/j.chemolab.2018.02.009>. URL <http://www.sciencedirect.com/science/article/pii/S0169743917304823>.
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RESEARCH PAPERS

A | A TOOL FOR SIMULATING MULTI-RESPONSE LINEAR MODEL DATA



A tool for simulating multi-response linear model data

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ABSTRACT

Data science is generating enormous amounts of data, and new and advanced analytical methods are constantly being developed to cope with the challenge of extracting information from such “big-data”. Researchers often use simulated data to assess and document the properties of these new methods, and in this paper we present an extension to the R-package simrel, which is a versatile and transparent tool for simulating linear model data with an extensive range of adjustable properties. The method is based on the concept of relevant components, and is equivalent to the newly developed envelope model. It is a multi-response extension of R-package simrel which is available in R-package repository CRAN, and as simrel the new approach is essentially based on random rotations of latent relevant components to obtain a predictor matrix X , but in addition we introduce random rotations of latent components spanning a response space in order to obtain a multivariate response matrix Y . The properties of the linear relation between X and Y are defined by a small set of input parameters which allow versatile and adjustable simulations. Sub-space rotations also allow for generating data suitable for testing variable selection methods in multi-response settings. The method is implemented as an update to the R-package simrel.

1. Introduction

Technological advancement has opened a door for complex and sophisticated scientific experiments that were not possible before. Due to this change, enormous amounts of raw data are generated which contain massive information but is difficult to excavate. Finding information and performing scientific research on these raw data has now become another problem. In order to tackle this situation new methods are being developed. However, before implementing any method, it is essential to test its performance and explore its properties. Often, researchers use simulated data for the purpose which itself is a time-consuming process. The main focus of this paper is to present a simulation method, along with an extension to the r-package called simrel, that is versatile in nature and yet simple to use.

The simulation method we are presenting here is based on the principle of relevant space for prediction [13] which assumes that there exists a y -relevant subspace in the complete space of predictor variables that is spanned by a subset of eigenvectors of these predictor variables. Our extension to this principle is to introduce a subspace in y (material space) which contains the information that predictor space is relevant for. The concept of response reduction to the material space in response variable was introduced by Cook et al. [6]. Our r-package based on this principle

lets the user specify various population properties such as; which latent components in x are relevant for a latent subspace of the responses y and the collinearity structure of x . This enables the possibility to construct data for evaluating estimation methods and methods developed for variable selection.

Among several publications on simulation, Johnson [16]; Ripley [17] and Gamerman and Lopes [9] have exhaustively discussed the topic. In particular, methods based on covariance structure has been discussed by Arteaga and Ferrer [2]; Arteaga and Ferrer [3] and Camacho [4], following approaches to find simulated data satisfying the desired correlation structure. In addition, many publications have implemented simulated data in order to investigate new estimation methods and prediction strategies [see:8, 5, 14]. However, most of the simulations in these studies were developed to address their specific problem. A systematic tool for simulating linear model data with single response, which could serve as a general tool for all such comparisons, was presented in Sæbø et al. [19] and as the r-package simrel. This paper extends simrel in order to simulate linear model data with multivariate response. The github repository of the package at <http://github.com/simulatr/simrel> has rich documentation with many examples and cases along with detailed descriptions of simulation parameters. In the following two sections, the discussion encircle the mathematical framework behind. In

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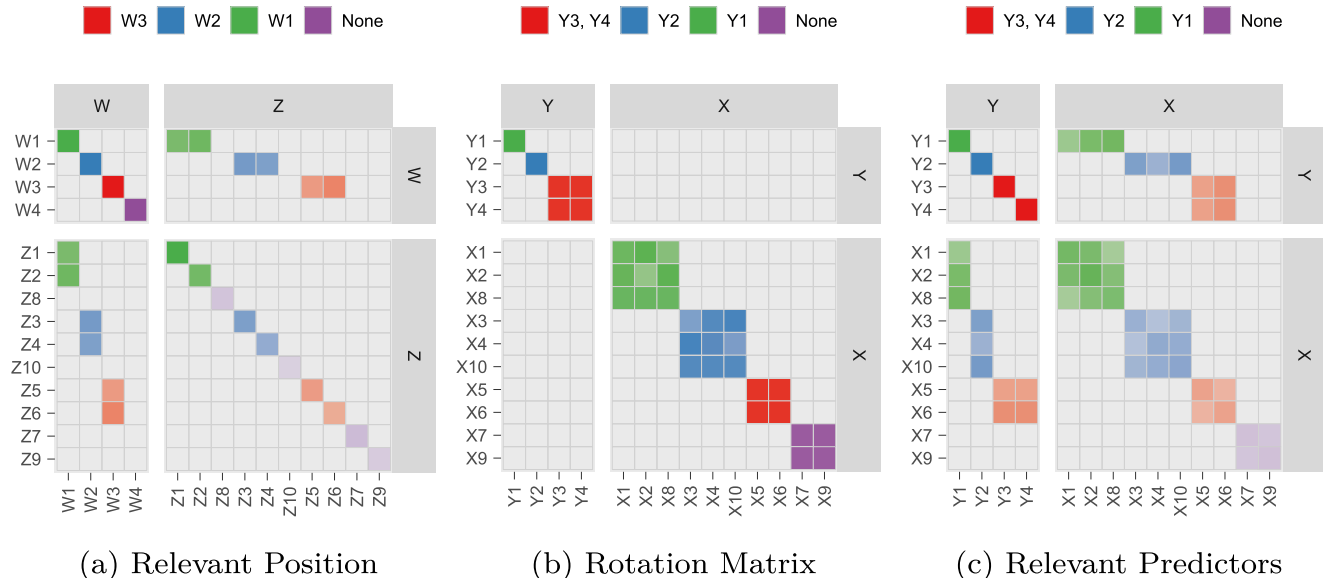


Fig. 1. Simulation of predictor and response variables after orthogonal transformation of predictor and response components by rotation matrices Q and R shown as the upper left and the lower right block matrices in (b).

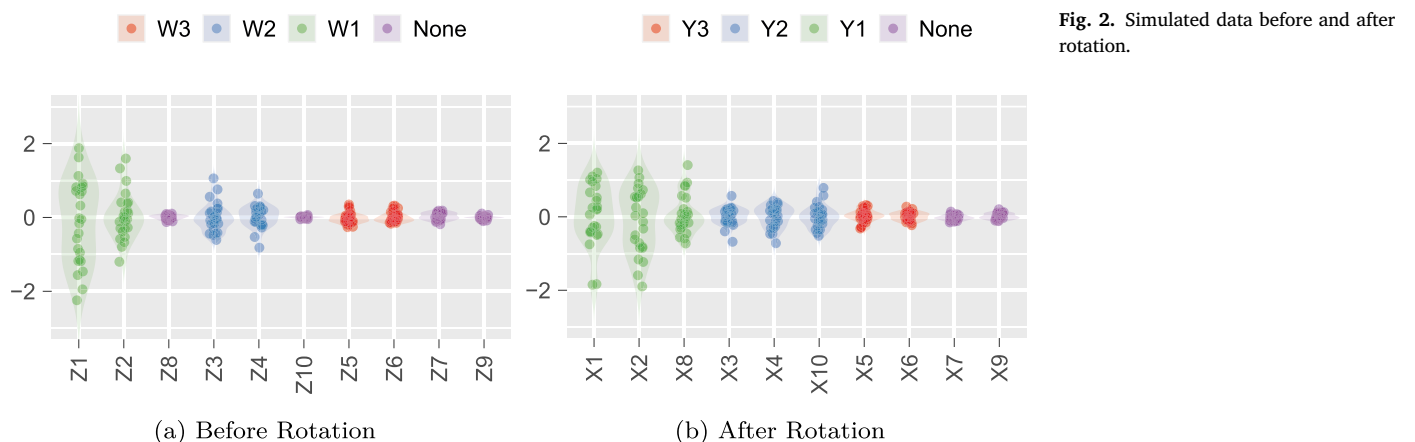


Table 1
Parameter setting of simulated data for comparison of estimation methods.

Decay of eigenvalues (γ)	Coef. of Determination ($\rho_{w_j}^2$)
Design1 0.2	0.8, 0.8, 0.4
Design2 0.8	0.8, 0.8, 0.4
Design3 0.2	0.4, 0.4, 0.4
Design4 0.8	0.4, 0.4, 0.4

addition, in section 4 and 5? we have also discussed the input parameters needed for simrel function in brief. In section 4, an implementation is presented as a case example and the final section introduces the shiny web application for this tool.

2. Statistical model

In this section we describe the model and the model parameterization which is assumed throughout this paper. We assume:

$$\begin{bmatrix} \mathbf{y} \\ \mathbf{x} \end{bmatrix} \sim N\left(\begin{bmatrix} \boldsymbol{\mu}_y \\ \boldsymbol{\mu}_x \end{bmatrix}, \begin{bmatrix} \Sigma_{yy} & \Sigma_{yx} \\ \Sigma_{xy}^T & \Sigma_{xx} \end{bmatrix}\right) \quad (1)$$

where, \mathbf{y} is a response vector with m response variables y_1, y_2, \dots, y_m with

mean vector $\boldsymbol{\mu}_y$, and \mathbf{x} is vector of p predictor variables with mean vector $\boldsymbol{\mu}_x$. Further,

Σ_{yy} ($m \times m$) is the variance-covariance matrix of \mathbf{y}
 Σ_{xx} ($p \times p$) is the variance-covariance matrix of variables \mathbf{x}
 Σ_{yx} ($m \times p$) is the matrix of covariance between \mathbf{x} and \mathbf{y}

Standard theory in multivariate statistics may be used to show that \mathbf{y} conditioned on \mathbf{x} corresponds to the linear model,

$$\mathbf{y} = \boldsymbol{\mu}_y + \boldsymbol{\beta}'(\mathbf{x} - \boldsymbol{\mu}_x) + \boldsymbol{\epsilon} \quad (2)$$

where, $\boldsymbol{\beta}'$ is a $(m \times p)$ matrix of regression coefficients, and $\boldsymbol{\epsilon}$ is an error term such that $\boldsymbol{\epsilon} \sim N(0, \Sigma_{y|x})$. The properties of the linear model (2) can be expressed in terms of covariance matrices in (1).

Regression Coefficients The matrix of regression coefficients is given by

$$\boldsymbol{\beta} = \boldsymbol{\Sigma}_{xx}^{-1} \boldsymbol{\Sigma}_{xy}$$

Coefficient of Determination Since, a matrix of coefficient-of-determination represents the proportion of variation explained by the predictors, we can write this matrix by its elements as,

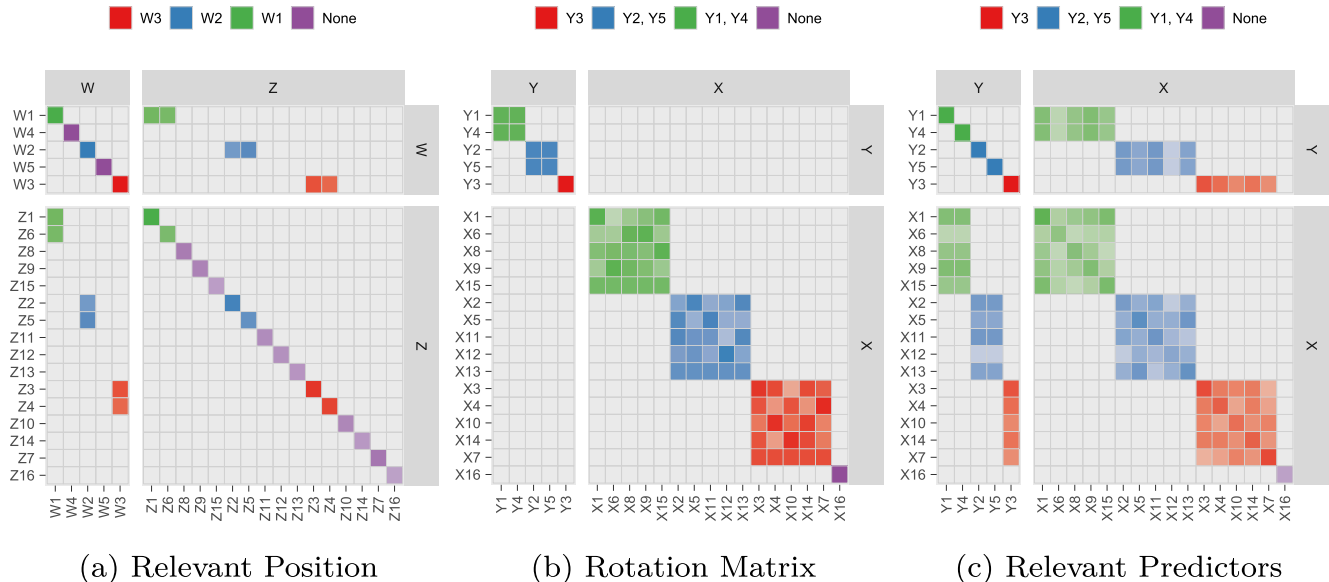


Fig. 3. Simulation of predictor and response variables for design one after orthogonal transformation of predictor and response components by rotation matrices Q and R shown as the upper left and the lower right block matrices in (b). Here (a) is the covariance structure of the latent space, which is rotated by the block diagonal rotation matrix in (b) resulting the covariance structure of simulated data in (c).

Table 2

Minimum average prediction error (number of components corresponding to minimum prediction error, minimum prediction error) (For Y_{env} , the number of response components is given).

Model	Design: 1	Design: 2	Design: 3	Design: 4
CPLS	(3, 3.24)	(4, 3.22)	(3, 4.09)	(3, 4.05)
CPPLS	(3, 3.21)	(3, 3.17)	(3, 4.11)	(3, 4.04)
OLS	(1, 3.60)	(1, 3.58)	(1, 4.57)	(1, 4.50)
PCR	(7, 3.28)	(6, 3.19)	(6, 4.08)	(6, 4.04)
PLS1	(2, 3.32)	(5, 3.20)	(1, 4.16)	(5, 4.07)
PLS2	(5, 3.29)	(6, 3.19)	(3, 4.11)	(6, 4.06)
Senv	(4, 3.17)	(5, 3.14)	(3, 4.35)	(5, 4.28)
Xenv	(5, 3.23)	(6, 3.20)	(5, 4.10)	(6, 4.11)
Yenv	(3, 3.24)	(3, 3.23)	(3, 4.29)	(3, 4.24)

$$\left(\rho_y^2\right)_{jj'} = \frac{\sigma_{xy}^t \Sigma_{xx}^{-1} \sigma_{xyj'}}{\sqrt{\sigma_{yj}^2 \sigma_{yj'}^2}} \forall j, j' = 1 \dots m$$

where, σ_{xyj} , $\sigma_{xyj'}$ are covariances between x and y_j , $y_{j'}$ respectively. Also, σ_{yj}^2 and $\sigma_{yj'}^2$ are unconditional variances of y_j and $y_{j'}$. Here the numerator is equivalent to the covariance of fitted y in sample space. if $j = j'$, it corresponds to a population version of the mean sum of squares of regression. The denominator gives the total unconditional variation in y . The diagonal elements of this matrix is the proportion of variation in a response $y_j, j = 1, \dots, m$ explained by the predictors.

Conditional variance The conditional variance-covariance matrix of y given x is,

$$\Sigma_{y|x} = \Sigma_{yy} - \Sigma_{yx} \Sigma_{xx}^{-1} \Sigma_{xy}.$$

The diagonal elements of this matrix equals the minimum least squared error of prediction $[E(y - \hat{y})^2]$ for each of the response variables.

Let us define a transformation of x and y as, $z = Rx$ and $w = Qy$. Here, $R_{p \times p}$ and $Q_{m \times m}$ are rotation matrices that rotate x and y to yield z and w , respectively. The model (1) can be re-expressed in terms of these transformed variables as:

$$\begin{aligned} \begin{bmatrix} w \\ z \end{bmatrix} &\sim N(\mu, \Sigma) = N\left(\begin{bmatrix} \mu_w \\ \mu_z \end{bmatrix}, \begin{bmatrix} \Sigma_{ww} & \Sigma_{wz} \\ \Sigma_{zw} & \Sigma_{zz} \end{bmatrix}\right) \\ &= N\left(\begin{bmatrix} Q\mu_y \\ R\mu_x \end{bmatrix}, \begin{bmatrix} Q\Sigma_{yy}Q' & Q\Sigma_{yx}R' \\ R\Sigma_{xy}Q' & R\Sigma_{xx}R' \end{bmatrix}\right) \end{aligned} \quad (3)$$

In addition, a linear model relating w conditioned on z can be written as,

$$w = \mu_w + \alpha'(z - \mu_z) + \tau \quad (4)$$

where α is the regression coefficient vector for the transformed model and $\tau \sim N(0, \Sigma_{w|z})$. Further, if both Q and R are orthonormal matrices, i.e., $Q^t Q = I_m$ and $R^t R = I_p$, the inverse transformation can be defined as,

$$\begin{aligned} \Sigma_{yy} &= Q\Sigma_{ww}Q' & \Sigma_{yx} &= Q'\Sigma_{wz} \\ \Sigma_{xy} &= R'\Sigma_{zw}Q & \Sigma_{xx} &= R'\Sigma_{zz}R \end{aligned} \quad (5)$$

From this, we can find a direct connection between different population properties of (2) and (4).

Regression Coefficients:

$$\alpha = \Sigma_{wz}^{-1} = Q\Sigma_{yz}R'[R\Sigma_{xx}R']^{-1} = Q[\Sigma_{yx}\Sigma_{xx}^{-1}]R' = Q\beta R'$$

Conditional Variance Further, the conditional variance-covariance matrix of w given z is,

$$\begin{aligned} \Sigma_{w|z} &= \Sigma_{ww} - \Sigma_{wz}\Sigma_{zz}^{-1}\Sigma_{zw} \\ &= Q\Sigma_{yy}Q' - Q\Sigma_{yx}R'[R\Sigma_{xx}R']^{-1}R\Sigma_{xy}Q' \\ &= Q\Sigma_{yy}Q' - Q\Sigma_{yx}\Sigma_{xx}^{-1}\Sigma_{xy}Q' \\ &= Q[\Sigma_{yy} - \Sigma_{yx}\Sigma_{xx}^{-1}\Sigma_{xy}]Q' = Q\Sigma_{y|x}Q' \end{aligned}$$

Coefficient of Determination The coefficient-of-determination matrix corresponding to w can be written as,

$$\begin{aligned} (\rho_w^2)_{jj'} &= \Sigma_{ww}^{-1/2} \Sigma_{wz} \Sigma_{zz}^{-1} \Sigma_{zw} \Sigma_{ww}^{-1/2} \\ &= \frac{\sigma_{zw}^t \Sigma_{zz}^{-1} \sigma_{zwj'}}{\sqrt{\sigma_{wj}^2 \sigma_{wj'}^2}} \forall j, j' = 1 \dots m \end{aligned}$$

where, σ_{zwj} and $\sigma_{zwj'}$ are covariances of z with w_j and $w_{j'}$, respectively. Also, σ_{wj}^2 and $\sigma_{wj'}^2$ are unconditional variances of w_j and $w_{j'}$. For simplicity,

Prediction Error

Averaged over 20 replicated Datasets of same properties

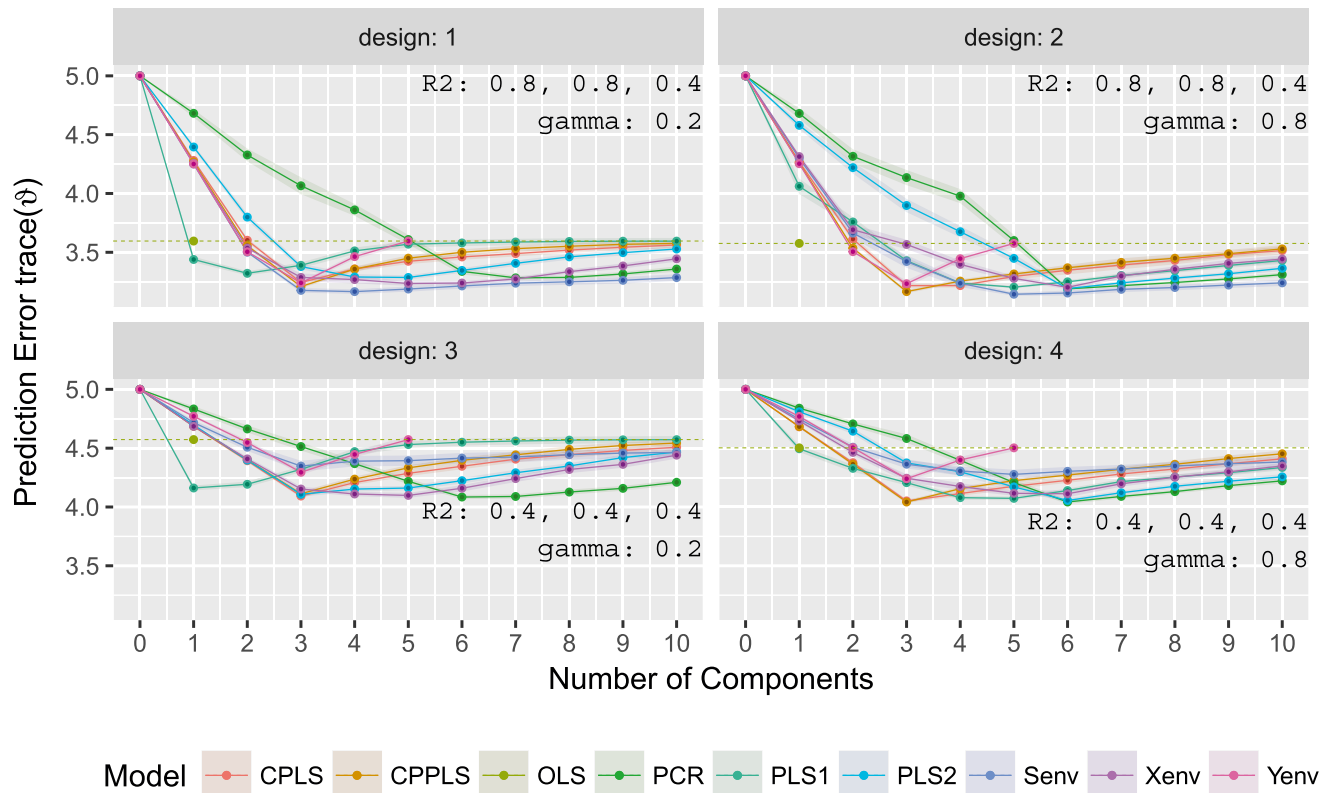


Fig. 4. Minimum of average prediction error.

Table 3
Simulation Design of second example.

$\eta : 0.1$	$\eta : 0.8$	Parameter	Value
Single Informative Response Component			
<i>Design 1</i>	<i>Design 2</i>	relpos	2, 3, 5, 7
		q	1000
		R2	0.8
Two Informative Response Components			
<i>Design 3</i>	<i>Design 4</i>	relpos	2; 3
		q	500; 500
		R2	0.6; 0.6

we will denote $\sigma_{z_i w_j}$ by σ_{ij} .

Since the rotation matrices give a direct connection between the covariance of (1) and (3), a straight forward relationship can be worked out between the terms in the above given matrix and their counterpart covariance matrices of the \mathbf{xy} -space.

From the eigenvalue decomposition principle, if $\Sigma_{xx} = \mathbf{RAR}^t$ and $\Sigma_{yy} = \mathbf{Q}\Omega\mathbf{Q}^t$ then \mathbf{z} and \mathbf{w} can be interpreted as principal components of \mathbf{x} and \mathbf{y} respectively. In this paper, these principal components will be termed as *predictor components* and *response components* respectively. Here, Λ and Ω are diagonal matrices of eigenvalues of Σ_{xx} and Σ_{yy} , respectively.

3. Relevant components

Consider a single response linear model with p predictors.

$$\mathbf{y} = \mu_y + \boldsymbol{\beta}'(\mathbf{x} - \mu_x) + \varepsilon$$

where, $\varepsilon \sim N(0, \sigma^2)$ and \mathbf{x} is a vector of random predictors. Following the concept of relevant space and irrelevant space which is discussed extensively in Helland and Almøy [13], Helland [12], Helland et al. [14], Cook et al. [5], and Sæbø et al. [19], we can assume that there exists a subspace of the full predictor space which is relevant for y . An orthogonal space to this space does not contain any information about y and is considered as irrelevant. Here, the y – relevant subspace of \mathbf{x} is spanned by a subset of the principal components defined by the eigenvectors of the covariance matrix of \mathbf{x} , i.e. Σ_{xx} .

This concept can be extended to m responses so that the subspace of \mathbf{x} is relevant for a subspace of \mathbf{y} . This corresponds to the concept of simultaneous envelopes [8] where relevant (material) and irrelevant (immaterial) space were discussed for both response and predictor variables.

3.1. Model parameterization

In order to construct a fully specified and unrestricted covariance matrix of \mathbf{z} and \mathbf{w} for the model in equation (3), we need to identify $1/2(p+m)(p+m+1)$ unknown parameters. For the purpose of simulation, we implement some assumptions to re-parameterize and simplify the model. This enables us to construct a wide range of model properties from only few key parameters.

Parameterization of Σ_{zz} If we let the rotation matrix \mathbf{R} correspond to the eigenvectors of Σ_{xx} , then \mathbf{z} becomes the set of principal components of \mathbf{x} . In that case Σ_{zz} is a diagonal matrix with eigenvalues $\lambda_1, \dots, \lambda_p$. Further, we adopt the same parametric representation as Sæbø et al. [19] for these eigenvalues:

$$\lambda_i = e^{-\gamma(i-1)}, \gamma > 0 \text{ and } i = 1, 2, \dots, p \quad (6)$$

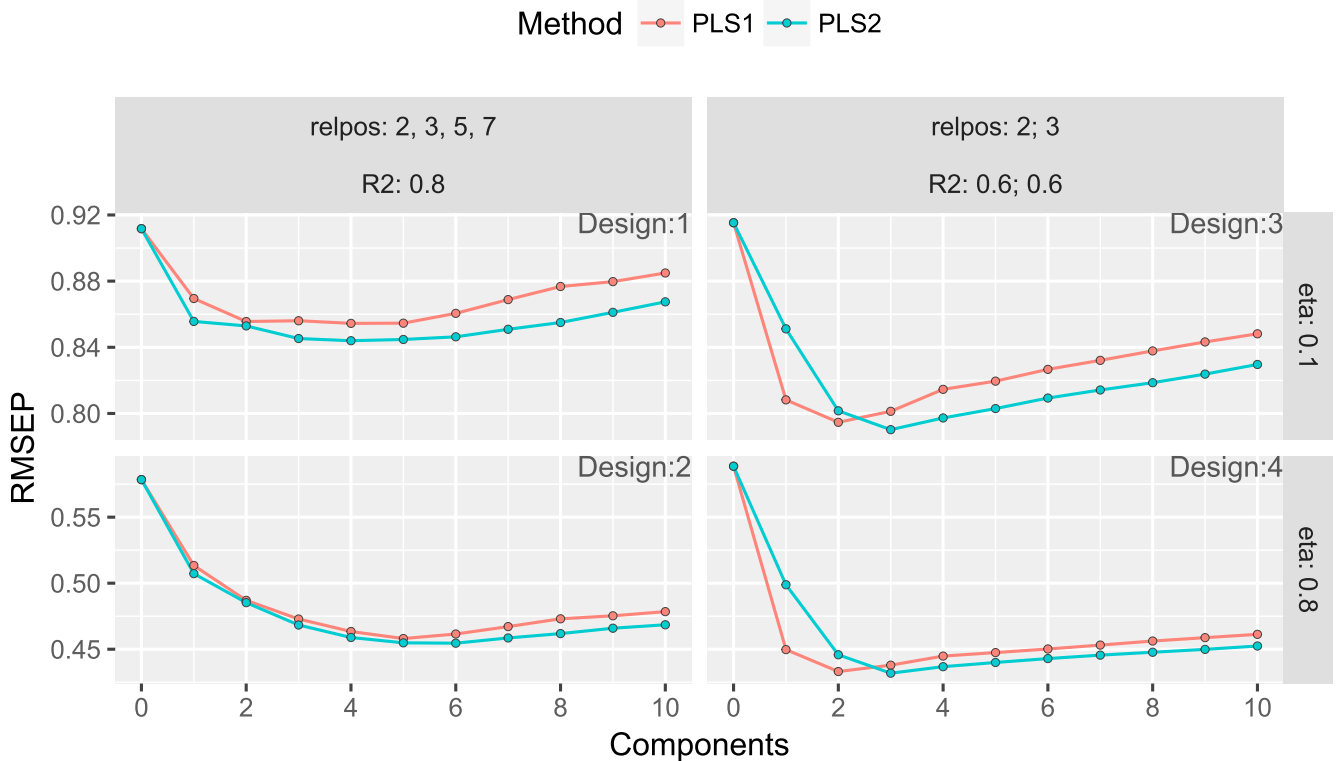


Fig. 5. Root mean square of error of prediction averaged over all response variables.

Here, as γ increases, the decline of eigenvalues becomes steeper, hence the parameter γ controls the level of multicollinearity in \mathbf{x} . We can write $\Sigma_{zz} = \Lambda = \text{diag}(\lambda_1, \dots, \lambda_p)$.

Parameterization of Σ_{ww} In similar manner, a parametric representation of eigenvalues corresponding to Σ_{ww} is adopted as,

$$\kappa_j = e^{-\eta(j-1)}, \eta > 0 \text{ and } j = 1, 2, \dots, m \quad (7)$$

Here, the decline of eigenvalues becomes steeper as η increases from zero. At $\eta = 0$, all w will have equal variance 1. Hence we can write $\Sigma_{ww} = \text{diag}(\kappa_1, \dots, \kappa_m)$.

Parameterization of Σ_{zw} After parameterization of Σ_{zz} and Σ_{ww} , we are left with $m \times p$ number of unknowns corresponding to Σ_{zw} . Some of the elements of Σ_{zw} may be equal to zero, which implies that the given z is irrelevant for the given variable w . The non-zero elements define which of the z that are relevant for w . We typically refer to the indices of these z variables as the positions of relevant components. In order to re-parameterize this covariance matrix, it is necessary to discuss the position of relevant components in detail.

3.1.1. Position of relevant components

Let k_1 components be relevant for w_1 , k_2 components be relevant for w_2 and so on. Let the positions of these components be given by the index sets $\mathcal{P}_1, \mathcal{P}_2, \dots, \mathcal{P}_m$ respectively. Further, the covariance between w_j and z_i is non-zero only if z_i is relevant for w_j . If σ_{ij} is the covariance between w_j and z_i then $\sigma_{ij} \neq 0$ if $i \in \mathcal{P}_j$ where $i = 1, \dots, p$ and $j = 1, \dots, m$ and $\sigma_{ij} = 0$ otherwise.

In addition, the true regression coefficients α for w_j (4) is given by:

$$\alpha_j = \Lambda^{-1} \sigma_{ij} = \sum_{i \in \mathcal{P}_j} \frac{\sigma_{ij}}{\lambda_i}, \quad j = 1, 2, \dots, m$$

The positions of the relevant components have heavy impact on prediction. Helland and Almøy [13] have shown that if the relevant components have large eigenvalues (variances), which here implies small index values in \mathcal{P}_j , prediction of y from x is relatively easy and if the

eigenvalues (variances) of relevant components are small, the prediction becomes difficult, given that the coefficient of determination and other model parameters are held constant. For example, if the first and second components, z_1 and z_2 , are relevant for w_1 and fifth and sixth components, z_5 and z_6 , are relevant for w_2 , it is relatively easier to predict w_1 than w_2 , other properties being similar. This might be so, because the first and second principal components have larger variances than the fifth and sixth components.

Although the covariance matrix may depend on few relevant components, we can not choose these covariances freely since we also need to satisfy following two conditions:

- The covariance matrices Σ_{zz} , Σ_{ww} and Σ must be positive definite
- The covariance σ_{ij} must satisfy user defined coefficient of determination

We have the relation,

$$\begin{aligned} \rho_w^2 &= \Sigma_{ww}^{-1/2} \Sigma_{zw}^t \Sigma_{zz}^{-1} \Sigma_{zw} \Sigma_{ww}^{-1/2} \\ &= \frac{\sigma_{ij}^t \Lambda^{-1} \sigma_{ij'}}{\sqrt{\sigma_{jj'}^2 \sigma_{jj}^2}} \forall j, j' = 1 \dots m \end{aligned}$$

Applying our assumptions that, $\Sigma_{ww} = \text{diag}(\kappa_1, \dots, \kappa_m)$ (7) and $\Sigma_{zz} = \Lambda = \text{diag}(\lambda_1, \dots, \lambda_p)$ (6), we obtain,

$$\rho_w^2 = \Sigma_{ww}^{-1/2} \Sigma_{zw}^t \Lambda^{-1} \Sigma_{zw} \Sigma_{ww}^{-1/2} = \begin{bmatrix} \sum_{i=1}^p \frac{\sigma_{i1}^2}{\lambda_i \kappa_1} & \cdots & \sum_{i=1}^p \frac{\sigma_{i1} \sigma_{im}}{\lambda_i \sqrt{\kappa_1 \kappa_m}} \\ \vdots & \ddots & \vdots \\ \sum_{i=1}^p \frac{\sigma_{i1} \sigma_{im}}{\lambda_i \sqrt{\kappa_1 \kappa_m}} & \cdots & \sum_{i=1}^p \frac{\sigma_{ii}^2}{\lambda_i \kappa_m} \end{bmatrix}$$

Furthermore, we assume that there are no overlapping relevant components for any two w , i.e., $\mathcal{P}_j \cap \mathcal{P}_{j^*} = \emptyset$ or $\sigma_{ij} \sigma_{ij^*} = 0$ for $j \neq j^*$. The additional unknown parameters in the diagonal of ρ_w^2 should agree with

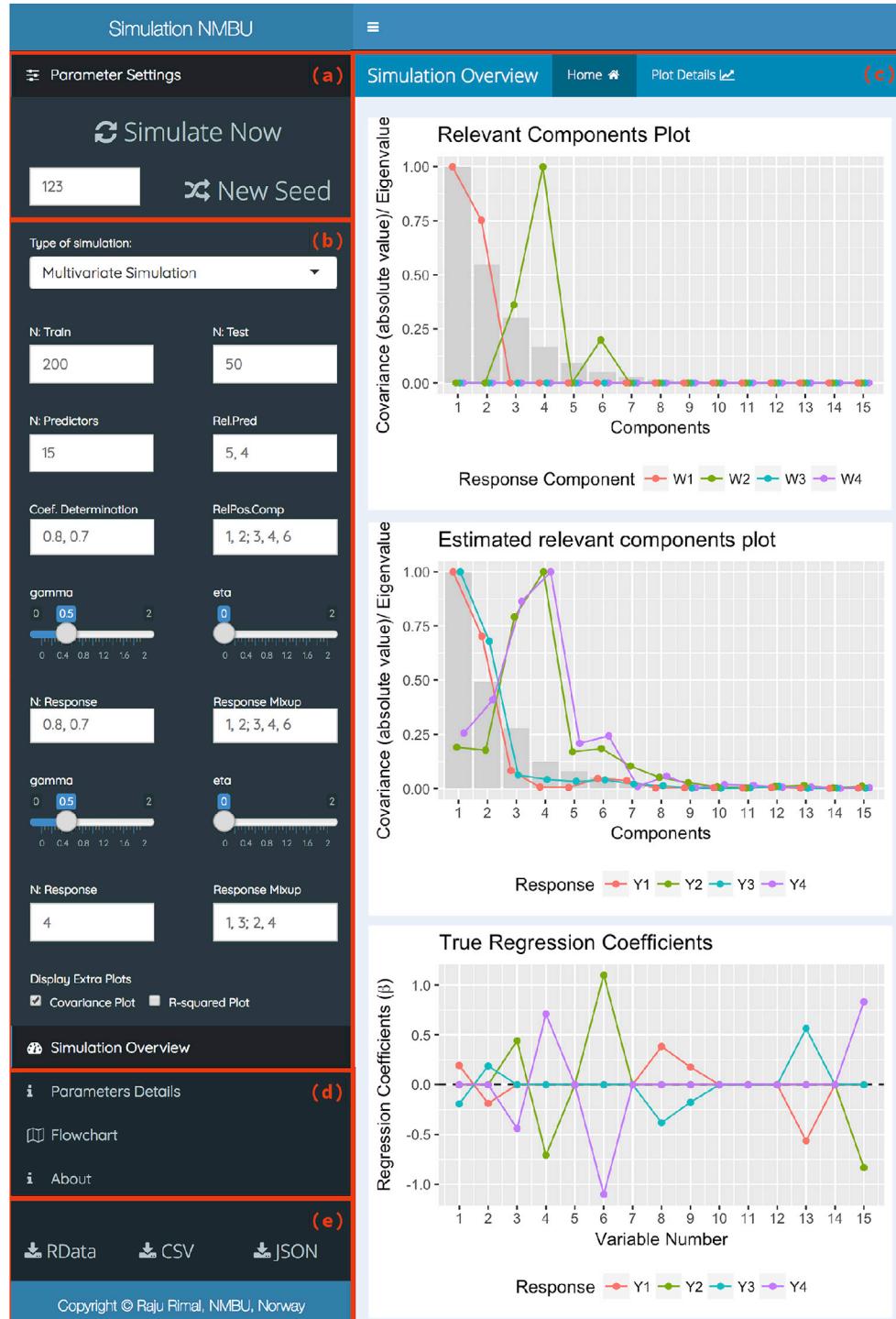


Fig. 6. Web interface of shiny application of simrel: (a) Buttons to trigger simulation, (b) Parameters for simulation, (c) Visualization of the true properties of simulated data (regression coefficients, true and estimated covariance between response and predictors components) (d) Additional analysis (e) Download option of simulated data.

user specified coefficients of determination for w . i.e., $\rho_{w_j}^2$ is,

$$\rho_{w_j}^2 = \sum_{i=1}^p \frac{\sigma_{ij}^2}{\lambda_i k_j}$$

Here, only the relevant components have non-zero covariances with w_j , so,

$$\rho_{w_j}^2 = \sum_{i \in \mathcal{P}_j} \frac{\sigma_{ij}^2}{\lambda_i k_j}$$

For some user defined $\rho_{w_j}^2$ the σ_{ij}^2 is determined as follows,

1. Sample k_j values from a uniform distribution $\mathcal{U}(-1, 1)$ distribution. Let them be denoted $\mathcal{S}_{\mathcal{P}_1}, \dots, \mathcal{S}_{\mathcal{P}_{k_j}}$.

2. Define,

$$\sigma_{ij} = \text{Sign}(\mathcal{S}_i) \sqrt{\frac{\rho_{w_j}^2 |\mathcal{S}_i|}{\sum_{k \in \mathcal{P}_j} |\mathcal{S}_k|}} \lambda_i k_j$$

for $i \in \mathcal{P}_j$ and $j = 1, \dots, m$

This means that the covariances between the predictor components and the response components are sampled randomly, but with restriction

```

simrel(
  n      = 200, # Number of training observations
  ntest  = 50, # Number of test observations
  p      = 15, # Number of predictor variables
  q      = c(5, 4), # Number of relevant predictors
  relpos = list(c(1, 2), c(3, 4, 6)),
  # Relevant predictor components
  R2    = c(0.8, 0.7), # Rsq for each response component
  m     = 4, # Number of response variables
  gamma = 0.6, # Decay factor of eigenvalues of predictors
  eta   = 0, # Decay factor of eigenvalues of responses
  ypos  = list(c(1, 3), c(2, 4)),
  # Combination of response components on rotation
  type  = "multivariate"
)

```

that the requested $\rho_{w_j}^2$ values are satisfied. This also implies that the regression coefficients α in (4) and β in (2) are sampled randomly under the same restriction.

3.1.2. Data simulation

From the above given parameterizations and the user defined choices of model parameters, a fully defined and known covariance matrix Σ of (\mathbf{w}, \mathbf{z}) is given. For the simulation of a single observation of (\mathbf{w}, \mathbf{z}) let us define $\mathbf{g} = \Sigma^{-1/2}\mathbf{u}$ such that $\text{cov}(\mathbf{g}) = \Sigma$. Here $\Sigma^{-1/2}$ is obtained from Cholesky decomposition of Σ , and \mathbf{u} is simulated from independent standard normal distribution.

Similarly, in order to simulate n observations, we define $\mathbf{G}_{n \times (m+p)} = \mathbf{U}\Sigma^{-1/2}$. Here the first m columns of \mathbf{G} will serve as \mathbf{W} and remaining p columns will serve as \mathbf{Z} . Further, each row of \mathbf{G} will be a vector sampled independently from the joint normal distribution of (\mathbf{w}, \mathbf{z}) . Finally, these simulated matrices \mathbf{W} and \mathbf{Z} are orthogonally rotated in order to obtain \mathbf{Y} and \mathbf{X} , respectively. In the following section we discuss these rotation matrices in more detail.

3.2. Rotation of predictor space

Initially, let us consider an example where a regression model with $p = 10$ predictors (\mathbf{x}) and $m = 4$ responses (\mathbf{y}). Let's assume that only three response components (w_1, w_2 and w_3) are needed to describe all four response variables. Further, let the index sets $\mathcal{P}_1 = \{1, 2\}$, $\mathcal{P}_2 = \{3, 4\}$ and $\mathcal{P}_3 = \{5, 6\}$ define the positions of the predictor components of \mathbf{x} that are relevant for w_1, w_2 and w_3 , respectively. Let $\mathcal{S}_1, \mathcal{S}_2$ and \mathcal{S}_3 be the orthogonal spaces spanned by each set of predictor components. These spaces together span $\mathcal{S}_k = \mathcal{S}_1 \oplus \mathcal{S}_2 \oplus \mathcal{S}_3$, which is the minimum relevant space and equivalent to the \mathbf{x} -envelope as discussed by Cook et al. [5].

Moreover, let $q_1 = 3, q_2 = 3$ and $q_3 = 2$ be the number of predictor variables we want to have relevant for w_1, w_2 and w_3 respectively. Then $q_1 = 3$ predictors may be obtained by rotating the predictor components in \mathcal{P}_1 along with one more irrelevant component. Similarly, $q_2 = 3$ predictors, relevant for w_2 , can be obtained by rotating predictor components in \mathcal{P}_2 along with one more irrelevant component and finally, $q_3 = 2$ predictors, relevant for w_3 , can be obtained by rotating the components in \mathcal{P}_3 without any additional irrelevant component. Let the

space spanned by the q_1, q_2 and q_3 number of predictors be $\mathcal{S}_{q_1}, \mathcal{S}_{q_2}$ and \mathcal{S}_{q_3} . Together they span a space $\mathcal{S}_q = \mathcal{S}_{q_1} \oplus \mathcal{S}_{q_2} \oplus \mathcal{S}_{q_3}$. This space is bigger than \mathcal{S}_k since in the process two irrelevant components were included in the rotations. Here, \mathcal{S}_k is orthogonal to \mathcal{S}_{p-k} and \mathcal{S}_q is orthogonal to \mathcal{S}_{p-q} . Generally speaking, here we are splitting the complete variable space \mathcal{S}_p into two orthogonal spaces – \mathcal{S}_k relevant for \mathbf{w} and \mathcal{S}_{p-k} irrelevant for \mathbf{w} .

In the previous section, we discussed about the construction of a covariance matrix for the latent structure. Fig. 1(a) shows a similar structure resembling the example here. The three colors represent the relevance with the three latent response components (w_1, w_2 and w_3). Here we can see that z_1 and z_2 (first and second predictor components of \mathbf{x}) have non-zero covariance with w_1 (first latent component of response \mathbf{y}). In the similar manner other non-zero covariances are self-explanatory.

In order to simulate predictor variables (\mathbf{x}), we construct matrix \mathbf{R} which then is used for orthogonal rotation of the predictor components \mathbf{z} . This defines a new basis for the same space as is spanned by the predictor components. In principle, there are many possible options for defining a rotation matrix. Among them, the eigenvector matrix of Σ_{xx} can be a candidate. However, in this reverse engineering approach both rotation matrices \mathbf{R} and \mathbf{Q} along with the covariance matrices Σ_{xx} are unknown. So, we are free to choose any \mathbf{R} that satisfies the properties of a real valued rotation matrix, i.e. $\mathbf{R}^{-1} = \mathbf{R}^t$ and $\det(\mathbf{R}) = \pm 1$ so that \mathbf{R} is orthonormal. Here the rotation matrix \mathbf{R} should be block diagonal as in Fig. 1(b) in order to rotate spaces $\mathcal{S}_1, \mathcal{S}_2, \dots$ separately. Fig. 2(a) shows the simulated predictor components \mathbf{z} that we are following in our example where we can see that the components z_1 and z_2 (relevant for w_1) is getting rotated together with an irrelevant component z_8 . The resultant predictors (Fig. 2(b)) x_1, x_2 and x_8 will hence also be relevant for w_1 . In the figure, we can see that components z_7, z_8, z_9 and z_{10} are not relevant for any responses before rotation, however, the x_8, x_{10} predictors become relevant after rotation keeping x_7 and x_9 still irrelevant.

Among several methods [1,11] for generating random orthogonal matrix, in this paper we are using orthogonal matrix \mathcal{Q} obtained from QR-decomposition of a matrix filled with standard normal variates. The rotation here can be a) restricted and b) unrestricted. The latter rotates all components \mathbf{z} together and makes all predictor variables somewhat relevant for all response components. However, the former performs a block-wise rotation so that it rotates certain selected predictor

components together. This gives control for specifying certain predictors as relevant for selected responses, which was discussed in our example above. This also allows us to simulate irrelevant predictors such as x_7 and x_9 which can be detected during variable selection procedures.

3.3. Rotation of response space

The previous example has four response variables with only three informative components w_1, w_2 and w_3 . During the rotation procedure, the response space is also rotated along with the predictor space. Fig. 1 shows that the informative response component w_3 is rotated together with the uninformative response component w_4 so that the predictors which were relevant for w_3 will be relevant for response variables y_3 and y_4 . Similarly, response components w_1 and w_2 are rotated separately so that predictors relevant for w_1 and w_2 will only be relevant for y_1 and y_2 respectively, which we can see in Fig. 2. Although the response components have exclusive set of relevant predictors, the rotation of the response space has the potential of creating several response variables that depend on the same relevant predictor space. In the r-package *simrel*, the combining of the response components is specified by a parameter *ypos*.

4. Implementation

This section demonstrates an application of multi-response extension of *simrel* with two examples in order to compare different estimation methods on the basis of prediction error. These examples are simply a demonstration of the use of *simrel* package rather than an extensive comparison of methods.

4.1. Example 1

For the comparison, we have considered four well established estimation methods.

- a) Ordinary Least Squares (OLS),
- b) Principal Component Regression (PCR),
- c) Partial Least Squares predicting individual response variable separately (PLS1) and
- d) Partial Least Squares predicting all response variables together (PLS2).

We have also considered four relatively new estimation methods in multi-response regression:

- a) Canonically Powered Partial Least Squares regression (CPPLS) [15],
- b) Canonical Partial Least Squares regression (CPLS) [15],
- c) Envelope estimation in predictor space (Xenv) [6],
- d) Envelope estimation in response space (Yenv) [7] and
- e) Simultaneous estimation of x- and y-envelope (Senv) [8].

From the possible combinations of two levels of coefficient of determination (ρ^2) and two levels of γ (6) (the factor that controls the multicollinearity in predictor variables), four simulation designs (design 1–4) were prepared. Replicating each design 20 times, 80 datasets with five response variables ($m = 5$) and 16 predictor variables ($p = 16$) were simulated using the method discussed in this paper. It was also assumed that three response components (w_1, w_2 and w_3) completely describe the variation present in five response variables ($y_1 \dots y_5$). Here, in this example we have assumed that all w 's have equal variance, i.e. $\Sigma_{ww} = I_m$, that is, $\eta = 0$ in (7). The four designs are presented in Table 1. All datasets contained 100 sampled observations and out of 16 predictor variables, three disjoint sets of five predictor variables each are relevant for response components w_1, w_2 and w_3 . Although the simulation method is well equipped to simulate data with $p \gg n$, for incorporating envelope estimation methods, which are based on maximization of likelihood, we

have chosen a $n > p$ situation in the example. Further, predictor components z_1 and z_6 were relevant for response component w_1 , predictor components z_2 and z_5 were relevant for response component w_2 and predictor component z_3 and z_4 were relevant for response component w_3 . In addition, following the discussion about rotation of response space (section 3.3), w_1 was rotated together with w_4 and w_2 was rotated together with w_5 . Fig. 3 visualizes the covariance structure and relationship between the response and predictor variables for the first design.

For each method, we can write an expected squared prediction error as,

$$\vartheta_{m \times m} = E \left[(\hat{\beta} - \beta)^t \Sigma_{xx} (\hat{\beta} - \beta) \right] + \Sigma_{y|x} \quad (8)$$

where, $\hat{\beta}$ is an estimate of the true regression coefficient β and Σ_{xx} is the true covariance structure of the predictor variables obtained from *simrel*. Also, $\Sigma_{y|x}$ is the true minimum error of the model. Here $\hat{\beta}$ varies across different estimation methods while the remaining terms are the same for each dataset design. The expression in (8) is estimated from 20 replicated calibration sets. Further, an overall prediction error of all responses is measured by the trace of ϑ (8).

The minimum prediction error (measured as discussed above) for nine estimation methods averaged over 20 replications of four designs are shown in Table 2. The table also gives the number of predictor components (response components in case of Yenv), a method has used in order to obtain the minimum of average prediction error.

Table 2 shows that the simultaneous envelope has prediction error of 3.17 and 3.14 in design 1 (with 4 components) and design 2 (with 5 components), respectively, which is smaller than other methods. However, the method was not able to show the same performance in design 3 and design 4. The PCR model has the smallest prediction error (4.08) from 6 components in design 3 and Canonically Powered PLS has minimum prediction error (4.04) from 3 components in design 4. In design 3, we can also see that the Canonical PLS method has second best performance with only three components. The number of components vary across different replicated dataset, but the component corresponding to minimum prediction error is discussed here. A detailed picture of prediction error for each estimation method obtained for each additional component is shown in Fig. 4. Although designs 2 and 4 have higher levels of multicollinearity, the performance of the estimation methods is indifferent to its effect. Since all methods, except OLS, are based on shrinking of estimates, they are less influenced by the multicollinearity problem.

The analysis presented in Fig. 4 has addressed some questions such as how methods work when there exist a true reduced dimension in response space, but also raised other questions like why they perform differently. For example, what is the reason for the decreasing relative performance of the simultaneous envelope method as the ρ^2 values are reduced? Does this depend on the dimensions and shape of the y envelopes? Since the example is merely intended as a demonstration of how *simrel* can be used in scientific study, a more elaborative studies would be necessary to answer such questions, but for this purpose *simrel* would be a powerful tool.

4.2. Example 2

In this second example, wide matrices with 100 observations and 1000 predictor variables were simulated. Since wide matrices are common in various fields such as genomics, spectroscopy and chemometrics, we set up this second example to compare two variants of partial least square regression – PLS1 and PLS2. While estimating regression coefficients PLS1 uses each response variable separately, while PLS2 uses them all simultaneously. A simulation design was constructed as in Table 3. With each design, 20 replicated datasets were simulated having five response variables and a moderate level of multicollinearity within the predictor variables ($\gamma = 0.5$).

The comparison were based on the prediction error measured by root mean square error of prediction (RMSEP). In order to approximate the error to theoretically computed error, 1000 extra test samples were drawn from the same distribution as the training samples during simulation.

One to ten components were used to fit the simulated data models. The prediction error was recorded for each response variable and each additional component. The first and second design in Table 3 has one informative response component for which four predictor components are relevant at positions 2, 3, 5 and 7, and the coefficient of determination is 0.8. Since the informative response component is rotated together with four uninformative response components, the information is shared among all five response variables after rotation.

The third and fourth design has two informative response components. The first response component has one relevant predictor component at position 2 and a coefficient of determination of 0.6. Similarly, the second response component has one relevant predictor component at position 3 and also here the coefficient of determination is 0.6.

In addition to having one and two response component models, two levels of variance structure of the response components is considered and defined by η parameters with values 0.1 and 0.8 respectively. In the first and third design, all response components vary in similar manner ($\eta = 0.1$), while in the second and fourth design the informative response components have higher variance ($\eta = 0.8$) than the uninformative ones as the eigenvalues of Σ_{yy} drop faster in this case.

Fig. 5 shows the average prediction error of test observations modelled by PLS1 and PLS2 for all four designs. The prediction errors are averaged over all 20 replicated datasets.

In general, PLS2 dominates PLS1 with regard to minimum error achieved for these simulated designs. The difference is largest for the designs with $\eta = 0.1$ in which case the response are moderately correlated and prediction appears to be more difficult than for $\eta = 0.8$. The effect of number of relevant response and predictor components appears to have less influence on the results than the covariance structure of Σ_{yy} . This small example of the use of simrel indicates that a more elaborate comparison study should be done on PLS1 and PLS2 in this respect.

5. Web interface

In order to give an alternative interface for simrel, we have created a shiny app which allows users to provide the simulation parameters through different input fields. Fig. 6 shows a screenshot of the application. The application contains three main sections through which the user can interact with this simulation approach. A random seed can be selected using section Fig. 6 (a) so that a particular set of data can be resimulated if needed. Fig. 6 (b) has all the input panels where the user-dependent parameters for simulation can be entered. Here the user also has the option to simulate univariate, bivariate or multivariate response data. In addition, a simulated R-object comprising the simulated data can be downloaded in Rdata format (section (e) in Fig. 6). The object holds the simulated data along with other properties such as coefficient of determination for each response, true regression coefficients and rotation matrices. Users can also download simulated data in JSON and CSV format.

All simrel parameters can be entered using a simple user interface where vector elements are separated with comma (,) and list elements are separated with semicolon (;). For instance, the relevant position discussed in the implementation (section 4) of this paper can be entered as 1, 6; 2, 5; 3, 4 which is equivalent to R syntax list (c(1, 6), c(2, 5), c(3, 4)). An R expression equivalent to the input parameters as shown in Fig. 5(b) can be written as,

With the parameters for simulation in the screenshot (Fig. 6) 200 training sets (n) and 50 test sets (ntest) will be simulated with 15 predictor variables (p) and 4 response variables (m). The 4 response variables will have a true latent dimension of two, which is spanned by the relevant *response components*. The first response component is rotated

together with the third (irrelevant) response component and the second response component is rotated together with the fourth (irrelevant) response component as set in ypos. Out of 15 predictors, 5 will be relevant for the first response component and 4 will be relevant for the second response component, as set by q. The 5 predictor variables, that are relevant for the first response component, span the same space as the predictor components at position 1 and 2. Similarly, the 4 predictor variables that are relevant for the second response component, span the same space as the predictor components at position 3, 4 and 6 (relops). The coefficient of determination for the first and second response components are 0.8 and 0.7, respectively (R2). The eigenvalues of the predictor components decay exponentially by the factor of 0.6 (gamma), whereas the eigenvalues of response components are constant (but can be set to exponential decay) (eta).

The application not only allows users to simulate data, but also gives some insight into simulated data properties. Section (c) in Fig. 6 contains three plots – a) true regression coefficients b) relevant components and c) estimated relevant components. In the first plot (Fig. 6(c) top) we can see that predictor variables (1, 2, 8, 9 and 13) are relevant for the first and third response variables (red and blue line) by their non-zero coefficients, whereas predictor variables (3, 4, 6 and 15) are relevant for the second and fourth response variables (purple and green line). The second plot (Fig. 6(c) middle) shows the covariances between the response components and the predictor components along with the corresponding eigenvalues in the background (bar plot). In the plot the absolute value of the covariances after scaling with the largest covariance are shown. As in our parameter setting, the plot shows that the first (red line) and second (green line) predictor components have non-zero covariance with the first and third response components, and the fourth and sixth predictor components have non-zero covariance with the second response component. The third plot (Fig. 6(c) bottom) is the estimated covariances between the predictor components and the response variables, for the simulated data. Since the first and third response components are rotated together, in the plot, the covariance between the predictor components and the first and third response variables (red and blue line) are following similar patterns as the theoretical (6(c) middle). This also suggests that the predictor components which were relevant for the first response component, becomes relevant for the first and third response variables after rotation.

Along with these main sections, section (d) in the same figure contains additional analysis performed on the simulated data such as its estimation with different methods. This section is intended for educational purposes to show how changing the data properties influences the performances of different estimation and prediction methods. Beside this application, for Rstudio users, a gadget will be available after installing the r-package. This gadget provides an interface enabling users to input simulation parameters and access some of the properties.

Many scientific studies [8,14,18] are using simulated data in order to compare their findings with others or assess its properties. In many of these situations, a user-friendly and versatile simulation tool like simrel can play an important role. Gangsei et al. [10] and Sæbø et al. [19] are some examples where the univariate and bivariate form of simrel have been used for such purposes.

6. Conclusion

Whether comparing methods or assessing and understanding the properties of any method, tool or procedure; simulated data allows for controlled tests for researchers. However, researchers spend enormous amount of time creating such simulation tools so that they can obtain a particular nature of data. We believe that this tool along with the R-package and the easy-to-use shiny web interface will become an assistive tool for researchers in this respect.

Appendix A. Supplementary data

Supplementary data related to this article can be found at <https://doi.org/10.1016/j.chemolab.2018.02.009>.

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B | MODEL AND ESTIMATORS FOR PARTIAL LEAST SQUARES REGRESSION



Model and estimators for partial least squares regression

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Abstract

Partial least squares (PLS) regression has been a very popular method for prediction. The method can in a natural way be connected to a statistical model, which now has been extended and further developed in terms of an envelope model. Concentrating on the univariate case, several estimators of the regression vector in this model are defined, including the ordinary PLS estimator, the maximum likelihood envelope estimator, and a recently proposed Bayes PLS estimator. These are compared with respect to prediction error by systematic simulations. The simulations indicate that Bayes PLS performs well compared with the other methods.

KEYWORDS

Bayes PLS estimator, envelope model, partial least squares, partial least squares model, simulation

1 | INTRODUCTION

Supervised learning from multivariate data is a central problem area in applied statistics and also in chemometrics. Specifically, let our task be to predict a single variable y from a p -dimensional variable \mathbf{x} , having data on n units. From a statistical point of view, a large number of learning methods are discussed in Hastie et al,¹ mainly under the ordinary multiple regression model. In chemometrics, partial least squares (PLS) regression is the dominating method.

Partial least squares regression has had a vigorous development in the chemometric literature since it was proposed by Wold et al² and Martens and Næs.³ The method has been extended in several directions, and its applications have been expanded to an increasing number of fields, for instance, genomic data.⁴ Both these issues have been discussed in detail in a recent paper by Mehmood and Ahmed,⁵ where a wealth of further references may be found.

Sometimes, the issue is prediction, but very often, one also see interpretations of scoring, loading, and correlation plots; see, for instance, Martens and Martens.⁶ Such plots are not unfamiliar to statisticians in principal component connections, but they are much more used by the chemometric society, and many scientists find them informative. They are plots of the sample variants of the latent variables and parameters defined by (3), (4), and (5) below and, thus, involve consistent estimates of these quantities when $n \rightarrow \infty$ and probably also in the more general case $p/n \rightarrow 0$.

In the beginning, the PLS method was to some extent neglected or turned down by statisticians (an exception among others was Frank and Friedman⁷; see also Helland^{8,9}), but it is now included as a tool among other biased regression methods by applied statisticians. For a general discussion paper with contributions both from mathematical statisticians and chemometrists, see Sundberg.¹⁰

Indeed, there was a difference in culture between chemometrists and statisticians then, and this difference still exists to a large extent. A statement by Munck et al¹¹ illustrates this, as seen from one side: "If chemometrics in its historical development had been limited to follow current scientific (and statistical) theories there would have been minimal progress in its wide applications today."

Recently, the difference in culture was discussed in some detail by Martens.¹² On the one hand, Martens makes the point that the field of Chemometrics has a lot to learn from other disciplines—mathematics, statistics, and computer science.

Among other things, he says that it will not be enough to have efficient “black box” algorithms. On the other hand, he accuses statisticians in general for a predilection for “macho mathematics,” concluding in part that Chemometrics need more statistics but not more statisticians. In other parts of the paper, he talks about bridging the gap between the 2 disciplines, an effort that we whole heartedly support.

This difference in culture may in part be related to the concepts of creativity and rigor, qualities which to some extent may be called complementary. One could say that one culture puts more emphasis on creativity, the other on rigor. Of course, this is a huge simplification. First, there is a lot of creativity among statisticians, also mathematical statisticians. Secondly, one should emphasize that precise thinking also should influence practice. A case of point is the following: Chung and Keleş¹³ recently proved that the PLS regression vector is inconsistent when $p/n \rightarrow k > 0$ under a wide set of conditions. This result is probably not too well known among chemometrists; some may have a tendency to put much confidence in PLS regression when $p \sim n$ or $p > n$. It is to be emphasized that the inconsistency result in Chung and Keleş¹³ is only concerned with estimation of the regression vector. The mathematical properties of PLS as a “prediction” method when $p > n$ are largely unknown, from a statistical point of view. There is much positive empirical evidence among applied researchers on these properties, but statisticians have only started to attack this problem, since it from an analytic point of view is very difficult. In particular, see the very recent paper by Cook and Forzani,¹⁴ where asymptotic expansions allowing both n and p to be large are developed for PLS prediction with 1 component.

It is true that chemometrists have had a leading edge in the development of PLS and of certain multivariate methods, in particular, with respect to visualization etc, and they still are ahead of statisticians in this sense.

Accepting this, an important general question is what mathematical statisticians can contribute with in this development. There are relatively few papers by mathematical statisticians investigating statistical properties of the PLS regression method itself. There are however several investigations on the shrinkage properties of PLS; see Krämer¹⁵ and references there, and also Foschi¹⁶ with references. Garthwaite¹⁷ offered a simple interpretation of PLS. Stone and Brooks¹⁸ and Naik and Tsai¹⁹ discuss different generalizations of PLS; in the latter paper also, consistency of PLS is proved. In Stoica and Söderström,²⁰ an asymptotic formulae related to PLS is derived. Chun and Keleş²¹ extends consistency to the case $p/n \rightarrow 0$, introduces a sparse PLS algorithm, and compares methods by simulation. In Krämer and Sugiyama,²² the degrees of freedom of PLS regression is discussed, and this concept is used in model selection. See also references in this last paper.

In Helland and Almøy,²³ several predictors in the random x regression model were compared asymptotically as $n \rightarrow \infty$, including principal component regression (PCR) and sample PLS regression (see the next section). The conclusion was that PCR is best for very large irrelevant eigenvalues (excluded from the prediction equation), whereas PLS regression tends to be best for intermediate irrelevant eigenvalues. Because the difference is extremely small for small irrelevant eigenvalues, and because very large irrelevant eigenvalues seldom occur in practice (and if they do, they should be included in the prediction equation), it was concluded that PLS regression is the method of choice in many cases. An additional argument for PLS over PCR is that PLS involves only choosing the number of components, whereas PCR also entails deciding which of the components should be included in the prediction.

As already mentioned, Cook and Forzani¹⁴ give an asymptotic expansion of the prediction error in PLS regression, which also is informative when $p \rightarrow \infty$, but mainly limited to 1 component. Results with several components are also announced.

A vital aspect in the history of statistics is the interplay between model and estimators. Once a model is formulated, one can in principle think of several estimators in this model. A statistician will talk about a “hard” model in terms of probability distributions—at least in terms of a model equation and a statement of correlation between terms in this model. This is a concept that has had and has a great success in a number of disciplines and is at the very core of statistics as a science. Our goal in the present paper is to show that this concept can be applied—and is useful—also in connection to PLS. Specifically, our purposes are to

- stress that PLS as an algorithm can be connected to a unique statistical model (known since 1990);
- formulate 5 different ways to present this model (known in the statistical literature since 2013);
- argue that the simplest way to present the model is through the concept of relevant components—a reduction of the random x regression model;
- review briefly some statistical investigations related to PLS;
- ask if the PLS algorithm may be improved by modifying the weights;
- argue that once the model is presented, the comparison of different estimators in the model is relevant;
- present a systematic tool (`simrel`) for comparing estimators in the model with relevant components;
- present the maximum likelihood estimator in the model;

- present a Bayes estimator connected to the model;
- and compare the PLS algorithm, the maximum likelihood estimator, and the Bayes estimator in a systematic simulation study, mainly with near collinear data.

Thus, in the PLS model, one can certainly discuss other estimators than the usual PLS regression estimator, which can be seen as originating by replacing population (co)variances in the model by sample (co)variances. Two examples are the maximum likelihood estimator of Cook et al,²⁴ see also Cook et al^{25,26} and Cook and Zhang,²⁷ and the Bayesian estimator of Helland et al.²⁸ By simulation, both these estimators have performed well compared with PLS regression under certain conditions, but they have their disadvantages. The maximum likelihood estimator cannot be used in the case when the data matrix has rank less than p , and the Bayesian estimator requires heavy computations, in particular, when p is large.

To compare estimators, we make vital use of the recently developed simulation package *simrel*; see Sæø et al.²⁹ It is very important to have such a tool in an area where it is difficult to obtain results by purely analytical means.

We emphasize that this paper is based upon reduction of the *random x* regression model. When considering latent variables from PLS, and when considering near collinearity in the observed \mathbf{x} -variables, it is natural to treat these \mathbf{x} -variables as random. It is our philosophy that this is also the best way to look upon model reduction. On the other hand, in the context of prediction, one could argue that one should condition upon the \mathbf{x} -variables and consider them as fixed. A prominent paper on PLS regression, taking fixed \mathbf{x} -variables in the basic model, is Krämer and Sugiyama,²² where further references can be found.

In recent years, there has been a rapidly growing statistical literature on the envelope model—a model generalizing the PLS model. In addition to the maximum likelihood estimation paper mentioned above, the most important papers seem to be Cook and Zhang,³⁰ where simultaneous reduction in the \mathbf{x} - and \mathbf{y} -space is proposed, and Cook and Zhang,³¹ where extensions to other regression methods than linear regression are discussed. More references can be found in these papers.

Model reduction in regression models is discussed in general from the point of view of rotations in the x -space in Helland.³²

The plan of this paper is as follows: In Section 2, we formulate the model in 5 different ways, which can be shown to be equivalent. In Section 3, we define 4 different estimators in the model, including the recent Bayes PLS estimator of Helland et al.²⁸ In Section 4, we ask the question if the ordinary PLS estimator with m components can be improved by forcing the weight vector at step $m + 1$ to vanish; the answer turns out to be negative. In Section 5, we describe the simulations done for comparison of estimators with respect to prediction error, and in Section 6, we give the results of the simulations. In Section 7, we illustrate the methods on a real data set. Finally, Section 8 is a discussion section.

2 | THE MODEL: SEVERAL FORMULATIONS

Take as a point of departure the linear model

$$y = \mu_y + \beta'(\mathbf{x} - \boldsymbol{\mu}_x) + \epsilon, \quad (1)$$

where β and \mathbf{x} are p -dimensional and the random predictor \mathbf{x} has mean $\boldsymbol{\mu}_x$ and covariance matrix Σ_{xx} , for simplicity, assumed nonsingular here (this can be relaxed to assuming $\beta \in \text{span}(\Sigma_{xx})$ in the case where this matrix is singular; see Cook et al.²⁴ and also C below). Independently, ϵ is distributed with mean 0 and variance σ^2 . When doing prediction from this model for near collinear data, a model reduction may be called for. Throughout this paper, a definite m -dimensional model reduction, which may be formalized in several equivalent ways, will be used. When this model holds, we say that we have an envelope model or a PLS model of dimension m or that there are m relevant components for prediction in the model.

- A. Given a subspace S of R^p , let \mathbf{P}_S be the projection upon S , and let \mathbf{Q}_S be the projection orthogonal to S . For simplicity, discuss the case where $\boldsymbol{\mu}_x = \mathbf{0}$. Let now S be the smallest space such that (1) $\mathbf{Q}_S\mathbf{x}$ is uncorrelated with $\mathbf{P}_S\mathbf{x}$ and (2) y is uncorrelated with $\mathbf{Q}_S\mathbf{x}$ given $\mathbf{P}_S\mathbf{x}$. In this case, we may say that $\mathbf{Q}_S\mathbf{x}$ contains no linear information about y , neither directly nor through $\mathbf{P}_S\mathbf{x}$. Consider a reduction of the data to $\mathbf{P}_S\mathbf{x}$.
- B. Here is an algebraic characterization, which turns out to be equivalent. For a matrix \mathbf{M} , define \mathbf{MS} as the space of vectors \mathbf{Mz} , as \mathbf{z} runs through S , and let S^\perp be the space perpendicular to S . Let now S be the smallest space in R^p such that (1) both $\Sigma_{xx}S \subseteq S$ and $\Sigma_{xx}S^\perp \subseteq S^\perp$ and (2) $\text{span}(\beta) \subseteq S$. In this case, we say that S is the Σ_{xx} -envelope of $\text{span}(\beta)$. It can be shown Cook et al³³ that the envelope always exists as the smallest space with the stated properties.

C. The regression vector β can always be expanded in terms of the eigenvectors \mathbf{d}_i of Σ_{xx} :

$$\beta = \sum_{i=1}^p \gamma_i \mathbf{d}_i. \quad (2)$$

In general, when there are coinciding eigenvalues in Σ_{xx} , this expansion is not unique. However, assume that this sum can be reduced to exactly m nonzero terms: $\beta = \sum_{i=1}^m \gamma_i \mathbf{d}_i$, where the \mathbf{d}_i correspond to different eigenvalues of Σ_{xx} . We then say that there are m relevant components for prediction in the model. This reduction can be imagined to take place by 2 mechanisms: (1) Some of the γ_i 's are really zero, and (2) there are coinciding eigenvalues in Σ_{xx} . Then, one can rotate such that it is enough with 1 eigenvector for each eigenspace in the sum. In this approach, it is important that we only know that there are m nonzero terms in the sum, not which terms that are nonzero. For a closer discussion of this, see Næs and Helland³⁴ and Helland and Almøy.²³

D. Consider the population version of the well-known PLS algorithm: Take $\mathbf{e}_0 = \mathbf{x} - \mu_x$, $f_0 = y - \mu_y$, and for $a = 1, 2, \dots, m$ compute successively:

$$\mathbf{w}_a = \text{cov}(\mathbf{e}_{a-1}, f_{a-1}), \quad t_a = \mathbf{w}'_a \mathbf{e}_{a-1}, \quad (3)$$

$$\mathbf{p}_a = \text{cov}(\mathbf{e}_{a-1}, t_a)/\text{var}(t_a), \quad q_a = \text{cov}(f_{a-1}, t_a)/\text{var}(t_a), \quad (4)$$

$$\mathbf{e}_a = \mathbf{e}_{a-1} - \mathbf{p}_a t_a, \quad f_a = f_{a-1} - q_a t_a.$$

It can be proved⁹ and is important in this connection that under the reduced model C, this algorithm stops automatically after m steps when $m < p$: It stops because $\mathbf{w}_{m+1} = \text{cov}(\mathbf{e}_m, f_m) = 0$. After those m steps, we get the representations

$$\mathbf{x} = \mu_x + \mathbf{p}_1 t_1 + \dots + \mathbf{p}_m t_m + \mathbf{e}_m, \quad y = \mu_y + q_1 t_1 + \dots + q_m t_m + f_m \quad (5)$$

with the corresponding PLS population prediction

$$y_{m,PLS} = \mu_y + q_1 t_1 + \dots + q_m t_m = \mu_y + \beta'_{m,PLS} (\mathbf{x} - \mu_x).$$

Theorem 1. (Helland⁹ and Cook et al²⁴)

- (a) The 2 conditions A and B on the space S are equivalent.
- (b) The models formulated by C and D are equivalent.
- (c) When there are m relevant components for prediction, the envelope space S has dimension m , and S can be taken as $\text{span}(\mathbf{w}_1, \dots, \mathbf{w}_m) = \text{span}(\mathbf{d}_1, \dots, \mathbf{d}_m)$.
- (d) When the envelope space has dimension m , there are m relevant components for prediction.
- (e) In this case, we have $\beta_{m,PLS} = \beta$.

Proof. (a) is proved in Cook et al,²⁴ Proposition 1 and (b) in Helland.⁹, Theorem 2. Finally, (c)-(e) and the equivalence with E below are contained in Cook et al.²⁴, Proposition 5 \square

In this sense, all the model formulations (A-D) are equivalent; they describe the same reduced model. In Helland⁹ and Cook et al,²⁴ a fifth equivalent formulation in terms of a Krylov sequence is also given:

E. S is also spanned by the vectors $\sigma_{xy}, \Sigma_{xx}\sigma_{xy}, \dots, \Sigma_{xx}^{m-1}\sigma_{xy}$, and m is the smallest integer such that $\beta = \Sigma_{xx}^{-1}\sigma_{xy}$ belongs to S .

The simplest way to express the model reduction implied by PLS seems to be C. In analogy with the equivalence between A and B, this can also be expressed as a reduction of the x vector. Consider again the centered case $\mu_x = \mathbf{0}$. For details, see Næs.³⁴

C'. Let \mathbf{R} be a nonrandom pxm matrix of rank m . Normalize such that $\mathbf{R}'\mathbf{R} = \mathbf{I}$. There are m relevant components $\mathbf{R}'\mathbf{x}$ for predicting y if and only if \mathbf{R} can be found such that (a) $\beta \in \text{span}(\mathbf{R})$ and (b) $\text{span}(\mathbf{R})$ is spanned by eigenvectors of Σ_{xx} .

Being a reduced model that can be motivated in so many different ways, it is definitively of interest to find a good estimator of the regression vector β under this model.

3 | ESTIMATORS IN THE PLS/ENVELOPE MODEL

Now that the PLS model is introduced, we will start to look at estimators of the parameters in this model, in particular, estimators of β , which will give prediction. Of special interest is estimators that perform well in the case of near collinear data. Some estimators are already known from the literature.

- a. The ordinary PLS estimator can be introduced as follows: With data (\mathbf{X}, \mathbf{y}) , take initial values $\mathbf{E}_0 = \mathbf{X} - \bar{\mathbf{x}}\mathbf{1}'$ and $\mathbf{f}_0 = \mathbf{y} - \bar{\mathbf{y}}\mathbf{1}$. Run the population PLS algorithm for A steps with population (co)variances replaced by sample (co)variances. Ordinarily, A is found by cross-validation or by similar means. Note that from D in Section 2, the m -step PLS model is characterized by $\mathbf{w}_{m+1} = \text{cov}(\mathbf{e}_m, \mathbf{f}_m) = \mathbf{0}$. Theoretically, when $A = m$, we cannot expect the sample weights $\hat{\mathbf{w}}_{m+1}$ to be zero. However, since any continuous function of the sample covariances and variances is consistent for the same function of the population covariances and variances, since $\hat{\mathbf{w}}_{m+1}$ through the PLS algorithm is such a function and since $\mathbf{w}_{m+1} = \mathbf{0}$, we will have $\lim_{n \rightarrow \infty} \hat{\mathbf{w}}_{m+1} = \mathbf{0}$ almost surely.
- b. The sparse regression SPLS of Chun and Keles²¹: This requires 2 effective tuning parameters, and it also aims at variable selection. Sparse partial least squares (SPLS) seems to be better than ordinary PLS in certain cases, also when variable selection is not an issue.
- c. When $\mathbf{S} = (\mathbf{X} - \bar{\mathbf{x}}\mathbf{1}')(\mathbf{X} - \bar{\mathbf{x}}\mathbf{1})'$ has rank p , which specifically requires $n > p$, the maximum likelihood estimator of β under the multinormal envelope model was given in Cook et al.²⁴ This estimator is of course very useful, but it cannot be used for small n . Modifications of the maximum likelihood estimator, which cover also this case, were recently indicated in Cook et al.²⁵ That paper also gives a MATLAB toolbox for maximum likelihood estimation in the envelope model and in several generalizations of this model. A faster algorithm for maximum likelihood estimation is discussed in Cook and Zhang.²⁷ Even faster algorithms with modifications to small sample size $n < p$ are recently described in Cook and Zhang,³⁵ and an R-package was recently described by Cook et al.²⁶
- d. Under a specific rotation-invariant prior, the Bayes estimator of β under the model with m relevant components was given in Helland et al.²⁸ This estimator was shown to be close to the best equivariant estimator, but it requires heavy computation.

The estimation was performed by a Markov Chain Monte Carlo approach. Specifically, for given m , and for observed centered data \mathbf{y} and \mathbf{X} , the likelihood function is proportional to

$$f(\mathbf{y}, \mathbf{X} | \nu, \gamma, \mathbf{D}, \sigma^2) \propto (\sigma^2)^{-n/2} \exp \left(-\frac{1}{2\sigma^2} \left(\mathbf{y} - \mathbf{X} \sum_{i=1}^m \gamma_i \mathbf{d}_i \right)' \left(\mathbf{y} - \mathbf{X} \sum_{i=1}^m \gamma_i \mathbf{d}_i \right) \right) \\ \times \left(\prod_{i=1}^p \nu_i \right)^{-n/2} \prod_{j=1}^n \exp \left(-\frac{1}{2} \mathbf{x}'_j \left(\sum_{i=1}^p \frac{1}{\nu_i} \mathbf{d}_i \mathbf{d}'_i \right) \mathbf{x}_j \right), \quad (6)$$

where $\nu = [\nu_1, \dots, \nu_p]$ and $\mathbf{D} = [\mathbf{d}_1, \dots, \mathbf{d}_p]$ are the eigenvalues and the eigenvectors of the \mathbf{x} -covariance matrix Σ_{xx} and $\gamma = [\gamma_1, \dots, \gamma_m]$ are regression parameters of the PLS model.

As argued in Helland et al.,²⁸ a near optimal equivariant regressor is found as the Bayesian estimator under rotation-invariant prior for $\mathbf{d}_1, \dots, \mathbf{d}_p$ and prior $\pi(\gamma) = \prod_i 1/\gamma_i^{1-\epsilon}$, where $1/\epsilon$ is a large uneven integer. Slightly modified scale priors are also chosen for ν as $\pi(\nu) = \prod_i 1/\nu_i \exp(-\epsilon_\nu/2\nu_i)$ and for σ^2 as $\pi(\sigma^2) = 1/\sigma^2 \exp(-\epsilon_\sigma/2\sigma^2)$. Here, ϵ_ν and ϵ_σ are some small numbers chosen to ensure properness of the posterior distribution. Estimation of model parameters may be done by means of Markov chain Monte Carlo methods. As shown in Helland et al.,²⁸ the marginal posterior distributions for σ^2 and ν_i (for $i = 1, \dots, p$) are, for the given prior distributions, all inverse gamma distributions. Furthermore, the marginal posterior distributions for γ_i (for $i \in 1, \dots, m$) are approximately normally distributed. There is no closed form posterior distribution for \mathbf{D} , hence a random walk step with a Metropolis-Hastings acceptance step is necessary for the sampling from the posterior distributions of the parameters. R-code for the Bayes estimator is available at <http://www.github.com/solvs/BayesPLS>, and further details on the Markov chain Monte Carlo implementation may be found in the supplementary documentation to Helland et al.²⁸

By simulation, both the maximum likelihood estimator c and the Bayes estimator d were shown to perform well compared to the PLS estimator a. These 2 estimators assume a multinormal distribution of the data in their derivation, but the estimators themselves are valid under more general assumptions. Both the chemometric tradition and the envelope model of Cook et al^{24,33} demand no detailed distributional assumptions.

4 | CAN A BETTER ESTIMATOR BE FOUND BY SIMPLE MEANS?

The m step PLS model is characterized by the constraint $\mathbf{w}_{m+1} = \text{cov}(\mathbf{e}_m, f_m) = \mathbf{0}$. However, in the sample PLS algorithm, $\hat{\mathbf{w}}_{m+1}$ is a continuous random variable if the data are continuous. Hence, almost surely, $\hat{\mathbf{w}}_{m+1} \neq \mathbf{w}_{m+1} = \mathbf{0}$. This means that the estimator of the vector of PLS parameter falls outside the corresponding parameter space. On the other hand, by standard statistical theory, the maximum likelihood estimator and any Bayes estimator are always in the parameter space.

In this section, we ask the question whether we can improve the PLS algorithm in some way such that $\hat{\mathbf{w}}_{m+1} = \mathbf{0}$ for the improved algorithm. That is, we seek modified weights $\hat{\mathbf{w}}_1, \dots, \hat{\mathbf{w}}_m$ such that $\hat{\mathbf{w}}_{m+1} = \mathbf{0}$ in the modified algorithm. Unfortunately, the answer to this question is no. This programme is only possible when \mathbf{S} is invertible, and then it by necessity leads to the least squares solution. Let $\hat{\mathbf{W}}_A = (\hat{\mathbf{w}}_1, \dots, \hat{\mathbf{w}}_A)$ for any A .

First, we need some properties of the ordinary PLS algorithm.

Proposition 2. *At each step, the PLS weights satisfy*

$$\hat{\mathbf{w}}_{A+1} = \mathbf{s} - \hat{\mathbf{W}}_A' (\hat{\mathbf{W}}_A \hat{\mathbf{S}} \hat{\mathbf{W}}_A)^{-1} \hat{\mathbf{W}}_A' \mathbf{s}, \quad (7)$$

and the A step regression vector is

$$\hat{\beta}_A = \hat{\mathbf{W}}_A (\hat{\mathbf{W}}_A' \hat{\mathbf{S}} \hat{\mathbf{W}}_A)^{-1} \hat{\mathbf{W}}_A' \mathbf{s}. \quad (8)$$

Proof. These relations were proved in Helland,⁸ see equations (3.3) and (3.7) there, and were also used in Cook et al.²⁴ □

Now, fix m . To find an algorithm such that $\hat{\mathbf{w}}_{m+1} = \mathbf{0}$, we will have to modify the weights $\hat{\mathbf{w}}_1, \dots, \hat{\mathbf{w}}_m$.

Definition 1. For the purpose of this section, call a restricted PLS prediction any prediction method based on an estimator of β of the form (8) for $A = m$ such that

- 1.) $\hat{\mathbf{W}}_m = (\hat{\mathbf{w}}_1, \dots, \hat{\mathbf{w}}_m)$ is modified with respect to PLS in some way.
- 2.) Equation 7 holds for $A = m$ and gives $\hat{\mathbf{w}}_{m+1} = \mathbf{0}$.

Theorem 3. *An RPLS prediction method exists if and only if \mathbf{S} is invertible and $\mathbf{S}^{-1}\mathbf{s} \in \text{span}(\hat{\mathbf{W}}_m)$. In that case, $\hat{\beta}$ is equal to the least squares estimator $\mathbf{S}^{-1}\mathbf{s}$.*

Proof. Assume that (7) holds for $A = m$ and $\hat{\mathbf{w}}_{m+1} = \mathbf{0}$. Then, $\mathbf{s} = \hat{\mathbf{W}}_m (\hat{\mathbf{W}}_m' \hat{\mathbf{S}} \hat{\mathbf{W}}_m)^{-1} \hat{\mathbf{W}}_m' \mathbf{s}$. This is possible for general \mathbf{s} only if \mathbf{S} is nonsingular, and then it is equivalent to $\mathbf{R}\sqrt{\mathbf{S}^{-1}}\mathbf{s} = \sqrt{\mathbf{S}^{-1}}\mathbf{s}$ with $\mathbf{R} = \mathbf{A}(\mathbf{A}'\mathbf{A})^{-1}\mathbf{A}'$, where $\mathbf{A} = \sqrt{\mathbf{S}}\hat{\mathbf{W}}_m$. Since \mathbf{R} is the projector upon $\text{span}(\mathbf{A})$, this is again equivalent to $\sqrt{\mathbf{S}^{-1}}\mathbf{s} \in \text{span}(\sqrt{\mathbf{S}}\hat{\mathbf{W}}_m)$, or $\mathbf{S}^{-1}\mathbf{s} \in \text{span}(\hat{\mathbf{W}}_m)$. Then, putting $\mathbf{s} = \hat{\mathbf{W}}_m \mathbf{q}$ in (8) for some \mathbf{q} gives $\hat{\beta} = \hat{\mathbf{W}}_m \mathbf{q} = \mathbf{S}^{-1}\mathbf{s}$. □

Thus, Theorem 3 shows clearly that it is not possible to modify the PLS weights in a nontrivial way such that the modified estimator belongs to the parameter space.

5 | DATA SIMULATIONS FOR COMPARISON OF ESTIMATORS

A comparative study of the prediction performances of the regular PLS algorithm, the maximum likelihood envelope method, the Bayes PLS, and the method of ordinary least squares (OLS) was performed on data simulated from the random regression model (1) and a real dataset measuring various properties and near infrared (NIR) spectra of diesel fuels. This and the following section will focus on simulation study in detail. In the study, we consider envelope method for predictor reduction²⁴ and use R-code discussed in Cook et al.²⁶ A detailed description of the simulation procedure can be found in Sæbø et al²⁹ with the accompanying R-package simrel, but key features of the approach are presented next. The simulation set up is best explained from reexpressing model (1) in the Gaussian case as

$$\begin{bmatrix} y \\ \mathbf{x} \end{bmatrix} \sim \mathcal{N}(\boldsymbol{\mu}_{yx}, \Sigma_{yx}) = \mathcal{N}\left(\begin{bmatrix} \boldsymbol{\mu}_y \\ \boldsymbol{\mu}_x \end{bmatrix}, \begin{bmatrix} \sigma_y^2 & \boldsymbol{\sigma}_{xy}^t \\ \boldsymbol{\sigma}_{xy} & \Sigma_{xx} \end{bmatrix}\right), \quad (9)$$

where σ_{xy} is a vector holding the covariances between the predictors (\mathbf{x}) and the response (y). The vector of regression coefficients β is by standard theory given as $\beta = \Sigma_{xx}^{-1} \sigma_{yx}$, which in turn can be expressed in terms of the eigenvalues ν_1, \dots, ν_p and the eigenvectors $\mathbf{d}_1, \dots, \mathbf{d}_p$ of Σ_{xx} :

$$\beta = \sum_{i=1}^p \frac{\mathbf{d}_i' \sigma_{yx}}{\nu_i} \mathbf{d}_i = \sum_{i=1}^p \gamma_i \mathbf{d}_i, \quad (10)$$

as given in Equation 2. In `simrel`, the following simplifying assumptions are made:

- ▷ It is assumed that $\nu_i = e^{-\eta(i-1)}$ for $i = 1, \dots, p$, implying $\nu_1 = 1$ (which we may assume without loss of generality) and that all subsequent eigenvalues are decaying according to the size of the parameter η . A large η gives a rapid decrease in eigenvalues, implying high level of multicollinearity in \mathbf{x} .
- ▷ It is assumed that $m \leq p$ eigenvectors are relevant for y , which means that Equation 10 (potentially) reduces to

$$\beta = \sum_{i \in \mathcal{P}} \gamma_i \mathbf{d}_i, \quad (11)$$

where m -vector \mathcal{P} is the set of indices of the relevant components (relpos) for which $\gamma_i = \mathbf{d}_i' \sigma_{xy} / \nu_i \neq 0$. Hence, the envelope or the relevant space has dimension m (see Theorem 1).

- ▷ Without loss of generality, it is further assumed that $\sigma_y = 1$, $\mu_y = 0$ and $\mu_x = \mathbf{0}$.

In `simrel`, the actual values of σ_{xy} were set to satisfy a prespecified value of the population coefficient of determination ρ^2 . It may be shown that under the above assumptions, $\rho^2 = \sigma_{xy}' \Sigma_{xx}^{-1} \sigma_{xy}$. This completes the specification of the parameters used in `simrel`, and in the present comparison study, a design for the simulated data sets in terms of these parameters were as defined in Table 1.

From the possible combination of the above parameters, 32 calibration sets were simulated with 5 replications of each, ie, there were 160 calibration sets (datasets) altogether.

6 | SYSTEMATIC COMPARISONS

A systematic comparison of the methods across the simulation designs was made on the basis of their ability to predict test samples. Since the distribution of the simulated variables is fully known, the expected mean squared error of prediction (MSEP) based on some $\hat{\beta}$ estimated from a calibration set may be found as

$$E_x [E_y(y - \hat{y})^2] = \left[\sigma^2 + E(\hat{\beta} - \beta)^T \Sigma_{xx} (\hat{\beta} - \beta) \right] \frac{n+1}{n} \quad (12)$$

in the model. The expectation on the right-hand side of the above expression is estimated for each method and for each design as an average over the 5 replicated calibration sets. To study the effects of p , ρ^2 , relpos(\mathcal{P}), Method, and (η) along with their interactions, we first retrieved the minimum MSEP for each method across 1 to 10 components (assumed numbers of relevant components). In Figure 1, interaction plots for these data properties are displayed.

The effect of the third-order interaction between p , ρ^2 and Methods, which we see in Figure 1 (left), shows that the maximum likelihood-based estimation methods, in our case, the envelope and the OLS, perform poorly on data sets with large number of variables and low ρ^2 . Still, the performance of the envelope is better than OLS also in situations where $p = 40$ and $n = 50$, representing here $p \sim n$. The interaction plots suggest that the Bayes PLS and ordinary PLS estimation methods are better and more stable on average than the two other methods.

Similarly, the effect of third-order interaction between relpos, η , and Method in Figure 1 (right) shows that OLS method gives higher prediction error than other methods, but the effect of relpos is small but notable for the envelope method. Again, Bayes PLS and ordinary PLS are best.

TABLE 1 Parameters used for simulating calibration sets

Number of training samples	<i>n</i>	50
Number of predictor variables	<i>p</i>	15 and 40
Population coefficient of determination	ρ^2	0.5 and 0.9
Position of relevant components	\mathcal{P}	$\triangleright 1, 2 \triangleright 1, 3 \triangleright 2, 3$ and $\triangleright 1, 2, 3$
Decay factor of eigenvalues of Σ_{xx}	η	0.5 and 0.9

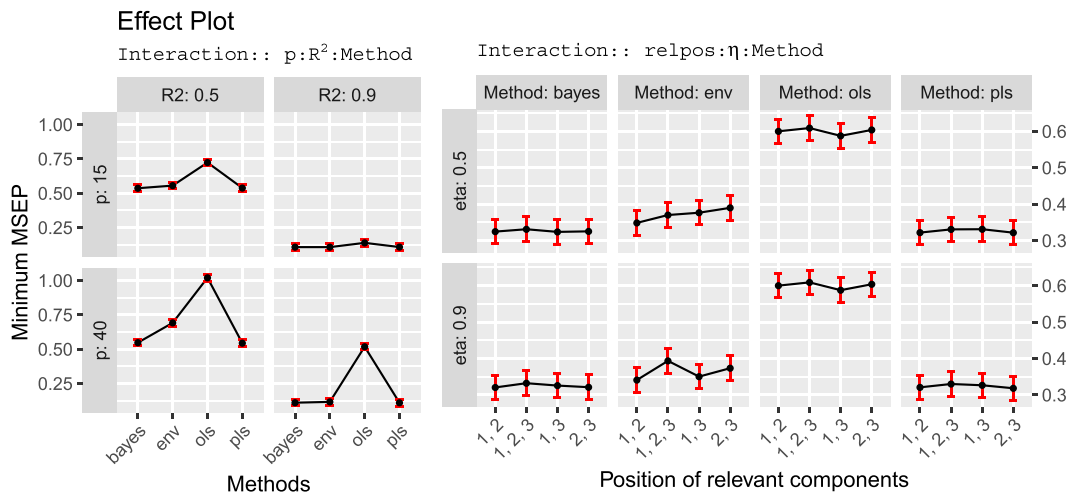


FIGURE 1 Third-order interaction effects. MSEP, mean squared error of prediction; ENV, envelope; OLS, ordinary least squares; PLS, partial least squares

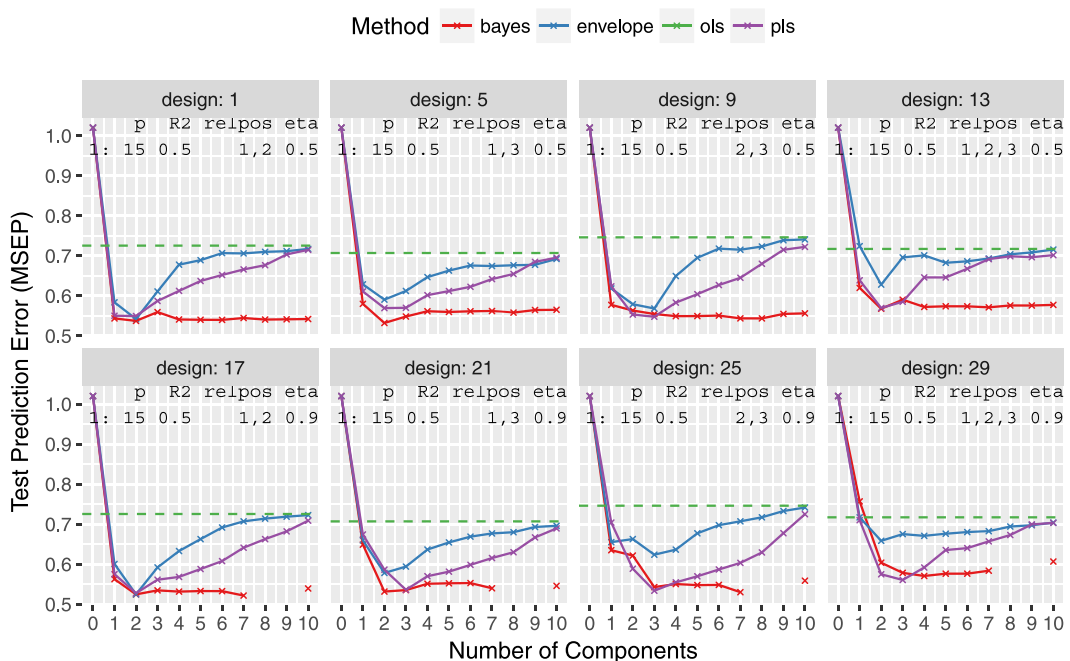


FIGURE 2 Average prediction error for designs with 15 predictor variables where coefficient of determination is 0.5. MSEP, mean squared error of prediction

The prediction error plots below are organized into 4 groups: (a) $p = 15$, $\rho^2 = 0.5$; (b) $p = 15$, $\rho^2 = 0.9$; (c) $p = 40$, $\rho^2 = 0.5$; and (d) $p = 40$, $\rho^2 = 0.9$. The OLS prediction error is shown by a straight dotted line.

In group (a), with small number of variables ($p \ll n$) and noisy data ($\rho^2 = 0.5$), Figure 2 shows that all the estimation methods performed better than OLS for all designs in this group, Bayes PLS being best in nearly all cases. Some convergence problems with Bayes PLS when eigenvalues decrease rapidly can be ignored since the minimum MSEP is already obtained from fewer components.

Having few variables rich with information ($\rho^2 = 0.9$), the designs in group (b) (Figure 3) leads to easy prediction with low prediction error in general for all methods. All the methods including OLS have small MSEPs, but the other methods are still dominant. In most of the situations, Bayes PLS has reached minimum error with only 1 component. In this group, the performance of envelope is better than regular PLS, and the minimum error for envelope is also achieved with fewer components.

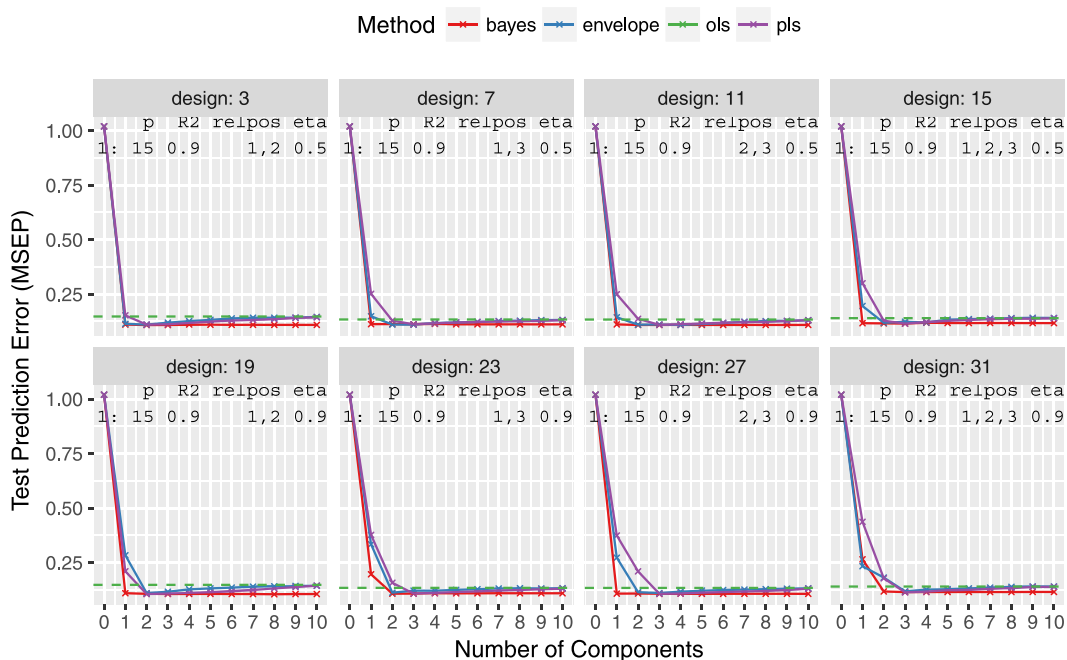


FIGURE 3 Average prediction error for designs with 15 predictor variables where coefficient of determination is 0.9. MSEP, mean squared error of prediction

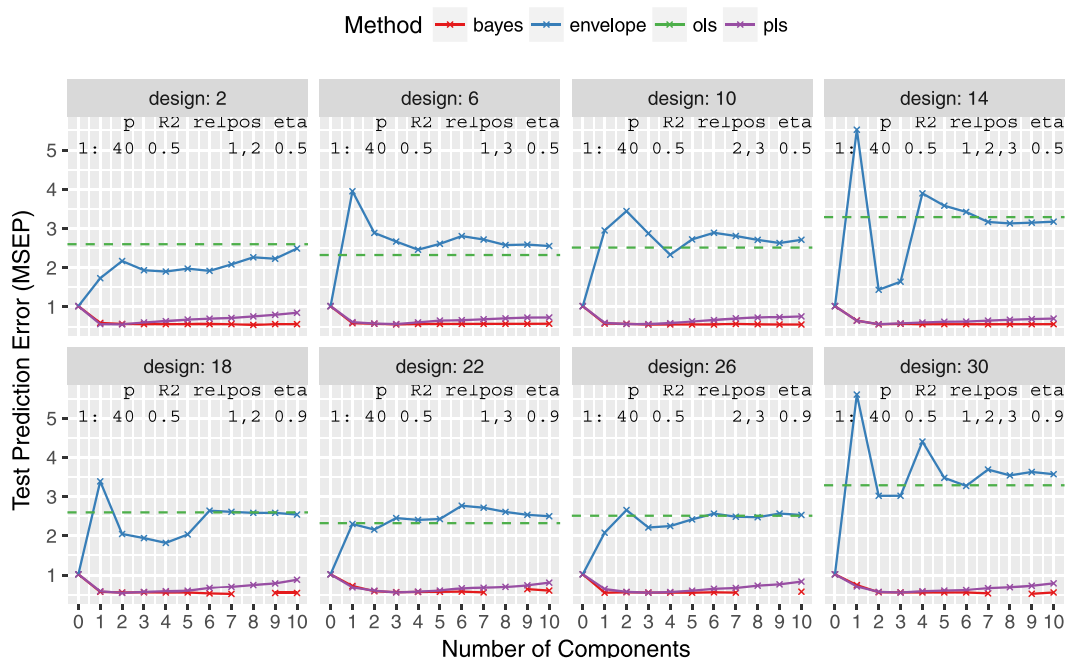


FIGURE 4 Average prediction error for designs with 40 predictor variables where coefficient of determination is 0.5. MSEP, mean squared error of prediction

Low information content combined with many predictor variables characterize the designs in group (c), and prediction is in general difficult for these designs. In Figure 4, the methods based on maximum likelihood estimation performed poorly and often poorer than an average guess. Bayes PLS and regular PLS performed well, as in the previous designs.

With 40 predictors ($p \sim n$) and rich information (high ρ^2) (designs in group d), Figure 5 shows that in most of the situations (except in design 16), the envelope method has nearly attained true minimum error (0.1) and has outperformed OLS. However, its prediction error is still larger than Bayes PLS and PLS. Bayes PLS and PLS methods are highly stable

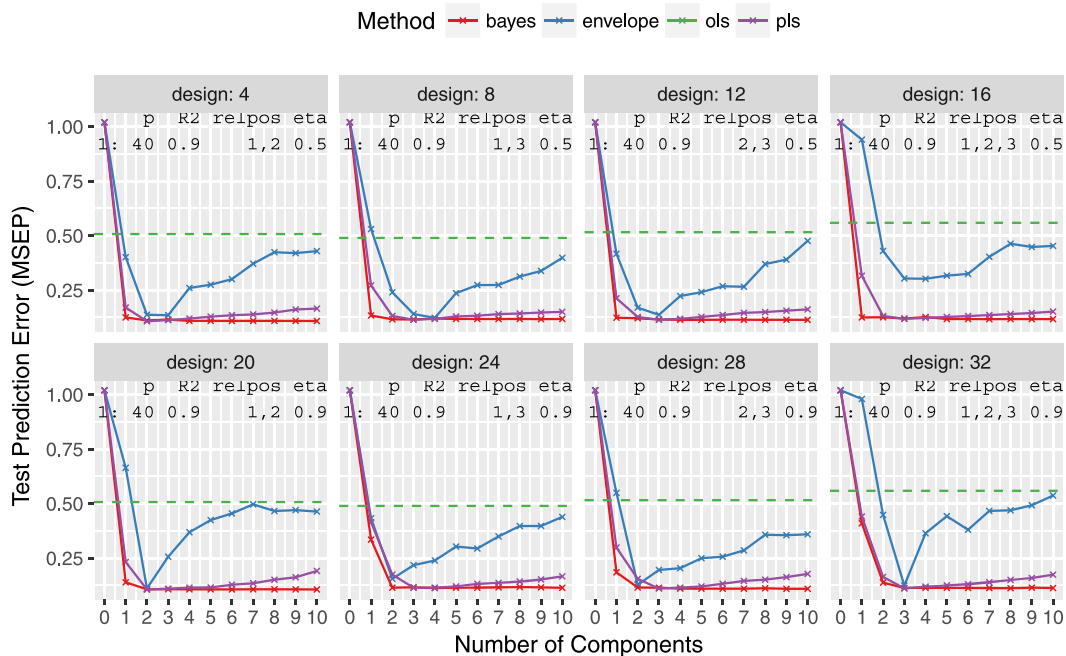


FIGURE 5 Average prediction error for designs with 40 predictor variables where coefficient of determination is 0.9. MSEP, mean squared error of prediction

and are closer to true minimum error. Further, Bayes PLS is able to obtain its minimum prediction error with only a small number of components.

In general, ordinary PLS is very stable in all situations. It is extensible (lots of variants have been developed after its introduction), easy, and less time consuming to fit than Bayes PLS and the envelope method. If the issue is to get closer prediction from squeezing information as much as possible, Bayes PLS will be a good alternative. Its performance with varying number of components is stable and better in all designs studied here. The envelope method performed better than OLS, and the performance increased for informative data ($\rho^2 = 0.9$). However, it has an increased error with additional components in many situations.

Correlation between estimated and true regression coefficients (β) along with the mean square error of estimation is presented for 4 designs in Figure 6. In case of ordinary PLS and the envelope method, the correlation for design 1 from group (a) and design 3 from group (b), both having 15 predictors, is high for small components. However, for design 2 from group (c) and design 4 from group (d), envelope methods exhibit sudden decrease in the correlation with corresponding increase in estimation error. The impressive prediction performance of Bayes PLS is also seen from the high correlation of estimated coefficients and true coefficients. In addition, the average mean square error of regression for this method is also small compared with others for all the components.

Although having low prediction error in case of envelope estimation method, the coefficient estimates are highly unstable for different components, which we can see from its variation in correlation with true coefficients (Figure 6, top). Bayes PLS and regular PLS estimates are more stable over different replicates and for different components (Figure 6, bottom) especially when $p \sim n$. This stability agrees with the low prediction error we have discussed before.

7 | COMPARISON OF ESTIMATORS USING NIR SPECTRA OF DIESEL FUELS

Let us consider an example using a real dataset. In this example, we have used data from <http://www.eigenvector.com/data/SWRI/>, which consists of NIR spectra of diesel fuels with different properties measured such as Catane Number. Since the variables in NIR spectra are highly correlated, we have selected a subset of every 10th variable as predictors and the property Catane Number as response. After removing missing observations, the first 150 observations were used as calibration set, and the rest 231 were used as validation set.

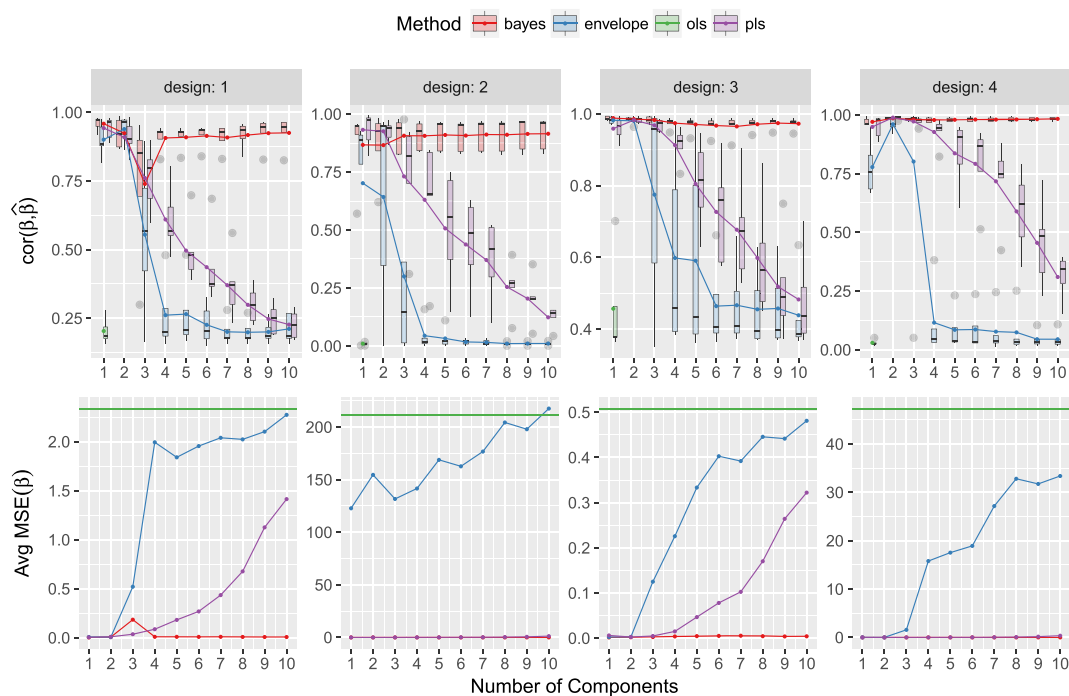


FIGURE 6 Correlation between true and estimated beta coefficient and beta estimation error. Box plots on the plots in first row show the variation in the correlation for each estimator and number of components used

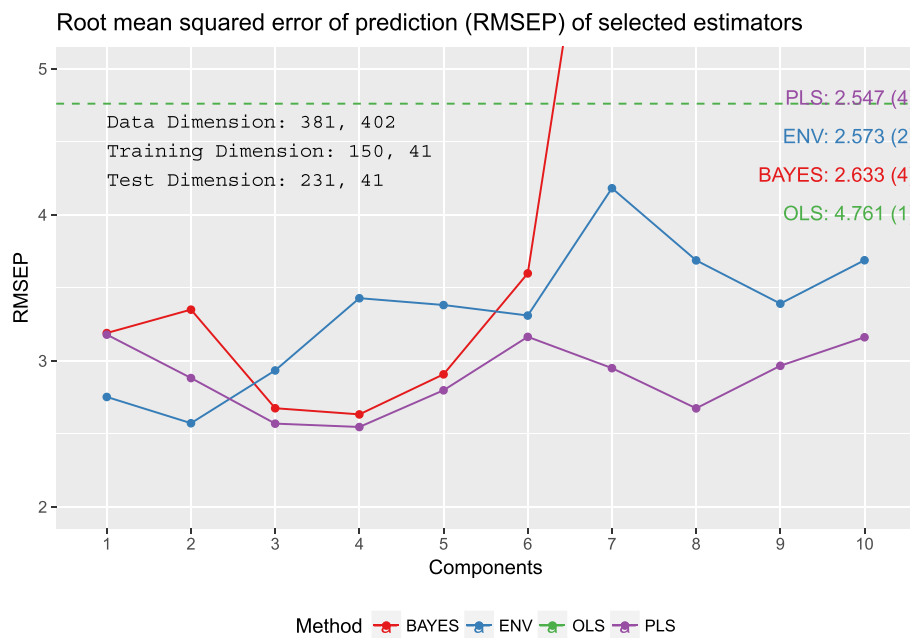


FIGURE 7 Root mean square error of prediction (RMSEP) from different estimators. Missing values were omitted in training and test datasets. ENV, envelope; OLS, ordinary least squares; PLS, partial least squares

Using the calibration set, a model with 1 to 10 components were fitted using PLS, envelope, and Bayes PLS methods. An OLS method was also fitted for reference. With each of these fitted models, the validation (test) set was used for prediction, and the root MSEP was measured. Based on the prediction error, Figure 7 compares the estimators we have considered.

The results from Figure 7 show quite different results from the systematic simulation study, mainly for Bayes PLS estimation. By using 3 and 4 components, the prediction from PLS and Bayes PLS is similar and can be considered their best. Envelope model is able to attain similar prediction error just in 2 components. It is important to notice that Bayes PLS

and envelope methods here are rather sensitive to the extra number of components, which also suggest that over-fitting must be examined before using the model for predicting new observations. In the example, all the methods have significant better performance than OLS.

8 | DISCUSSION

The purpose of the present article has been to discuss the approach to PLS regression via model reduction in the random \mathbf{x} multiple regression model, and to compare estimators in this reduced model.

From simulations, the Bayes estimator under the PLS model seems to have very good properties. In virtually all of the 32 designs, the MSEP curve for Bayes PLS lies below that for ordinary PLS and also that for the maximum likelihood envelope model. A particularly desirable feature of Bayes PLS is that the MSEP curve seems to be almost flat for varying number of components. Thus, the error made by choosing a wrong number of components m by cross-validation must be expected to be small.

Envelope and Bayes PLS estimation methods, when compared with ordinary PLS methods, display better prediction performance (only when p is small for the envelope method). However, both of them have their disadvantages. The envelope method, as based on maximum likelihood, breaks down when p approaches n , while Bayes PLS has time-consuming computation, and in our simulations, it failed to converge for some cases.

However, in the results in the example using real data, the performance of Bayes PLS estimator is in contrast to its result from the simulated data. Since the predictors are highly correlated, only a few number of components are sufficient for the prediction, but when an extra number of components were used, the estimators seem to be influenced by the noise which increases with each additional component. In this respect, a more thorough study on Bayes PLS should be done for its contrast results on simulated and real dataset. A convergence issue in Bayes PLS can be suspected for the reason as seen in the example using simulated data.

For practical purposes, the ordinary PLS algorithm still seems to be a good option for prediction purposes, but from a statistical point of view, a closer study of its properties as $p \rightarrow \infty$ seems to be called for. We feel that the model approach of the present paper may give a good framework for such a study, both in terms of asymptotic expansions and in terms of further simulations. Such simulations may also include the cross-validated LASSO and other methods such as ridge regression, but note that these estimators are derived from other considerations than that of predicting the effect of relevant components.

This paper has been concentrated on the case of univariate response. We hope to discuss the multivariate case later.

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C | COMPARISON OF MULTI-RESPONSE PREDICTION METHODS

Comparison of Multi-response Prediction Methods

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Abstract

While data science is battling to extract information from the enormous explosion of data, many estimators and algorithms are being developed for better prediction. Researchers and data scientists often introduce new methods and evaluate them based on various aspects of data. However, studies on the impact of/on a model with multiple response variables are limited. This study compares some newly-developed (envelope) and well-established (PLS, PCR) prediction methods based on real data and simulated data specifically designed by varying properties such as multicollinearity, the correlation between multiple responses and position of relevant principal components of predictors. This study aims to give some insight into these methods and help the researcher to understand and use them in further studies.

Keywords: model-comparison, multi-response, simrel

1. Introduction

The prediction has been an essential component of modern data science, whether in the discipline of statistical analysis or machine learning. Modern technology has facilitated a massive explosion of data however, such data often contain irrelevant information that consequently makes prediction difficult. Researchers are devising new methods and algorithms in order to extract information to create robust predictive models. Such models mostly contain predictor variables that are directly or indirectly correlated with other predictor variables. In addition, studies often consist of many response variables correlated with each other. These interlinked relationships influence any study, whether it is predictive modelling or inference.

Modern inter-disciplinary research fields such as chemometrics, econometrics and bioinformatics handle multi-response models extensively. This paper attempts to compare some multivariate prediction methods based on their prediction performance on linear model data with specific properties. The properties include the correlation between response variables, the correlation between predictor variables, number of predictor variables and the position of relevant predictor components. These properties are discussed more in the **Experimental Design** section. Among others, Sæbø et al. [25] and Almøy [1] have conducted a similar comparison in the single response setting. In addition, Rimal et al. [24] have also conducted a basic

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comparison of some prediction methods and their interaction with the data properties of a multi-response model. The main aim of this paper is to present a comprehensive comparison of contemporary prediction methods such as simultaneous envelope estimation (Senv) [7] and envelope estimation in predictor space (Xenv) [6] with customary prediction methods such as Principal Component Regression (PCR), Partial Least Squares Regression (PLS) using simulated dataset with controlled properties. In the case of PLS, we have used PLS1 which fits individual response separately and PLS2 which fits all the responses together. Experimental design and the methods under comparison are discussed further, followed by a brief discussion of the strategy behind the data simulation.

2. Simulation Model

Consider a model where the response vector (\mathbf{y}) with m elements and predictor vector (\mathbf{x}) with p elements follow a multivariate normal distribution as follows,

$$\begin{bmatrix} \mathbf{y} \\ \mathbf{x} \end{bmatrix} \sim \mathcal{N} \left(\begin{bmatrix} \boldsymbol{\mu}_y \\ \boldsymbol{\mu}_x \end{bmatrix}, \begin{bmatrix} \boldsymbol{\Sigma}_{yy} & \boldsymbol{\Sigma}_{yx} \\ \boldsymbol{\Sigma}_{xy} & \boldsymbol{\Sigma}_{xx} \end{bmatrix} \right) \quad (1)$$

where, $\boldsymbol{\Sigma}_{xx}$ and $\boldsymbol{\Sigma}_{yy}$ are the variance-covariance matrices of \mathbf{x} and \mathbf{y} , respectively, $\boldsymbol{\Sigma}_{xy}$ is the covariance between \mathbf{x} and \mathbf{y} and $\boldsymbol{\mu}_x$ and $\boldsymbol{\mu}_y$ are mean vectors of \mathbf{x} and \mathbf{y} , respectively. A linear model based on (1) is,

$$\mathbf{y} = \boldsymbol{\mu}_y + \boldsymbol{\beta}^t (\mathbf{x} - \boldsymbol{\mu}_x) + \boldsymbol{\epsilon} \quad (2)$$

where, $\boldsymbol{\beta}^t$ is a matrix of regression coefficients and $\boldsymbol{\epsilon}$ is an error term such that $\boldsymbol{\epsilon} \sim \mathcal{N}(0, \boldsymbol{\Sigma}_{y|x})$. Here, $\boldsymbol{\beta}^t = \boldsymbol{\Sigma}_{yx} \boldsymbol{\Sigma}_{xx}^{-1}$ and $\boldsymbol{\Sigma}_{y|x} = \boldsymbol{\Sigma}_{yy} - \boldsymbol{\Sigma}_{yx} \boldsymbol{\Sigma}_{xx}^{-1} \boldsymbol{\Sigma}_{xy}$

In a model like (2), we assume that the variation in response \mathbf{y} is partly explained by the predictor \mathbf{x} . However, in many situations, only a subspace of the predictor space is relevant for the variation in the response \mathbf{y} . This space can be referred to as the relevant space of \mathbf{x} and the rest as irrelevant space. In a similar way, for a certain model, we can assume that a subspace in the response space exists and contains the information that the relevant space in predictor can explain (Figure-1). Cook et al. [6] and Cook and Zhang [7] have referred to the relevant space as material space and the irrelevant space as immaterial space.

With an orthogonal transformation of \mathbf{y} and \mathbf{x} to latent variables \mathbf{w} and \mathbf{z} , respectively, by $\mathbf{w} = \mathbf{Q}\mathbf{y}$ and $\mathbf{z} = \mathbf{R}\mathbf{x}$, where \mathbf{Q} and \mathbf{R} are orthogonal rotation matrices, an equivalent model to (1) in terms of the latent variables can be written as,

$$\begin{bmatrix} \mathbf{w} \\ \mathbf{z} \end{bmatrix} \sim \mathcal{N} \left(\begin{bmatrix} \boldsymbol{\mu}_w \\ \boldsymbol{\mu}_z \end{bmatrix}, \begin{bmatrix} \boldsymbol{\Sigma}_{ww} & \boldsymbol{\Sigma}_{wz} \\ \boldsymbol{\Sigma}_{zw} & \boldsymbol{\Sigma}_{zz} \end{bmatrix} \right) \quad (3)$$

where, $\boldsymbol{\Sigma}_{ww}$ and $\boldsymbol{\Sigma}_{zz}$ are the variance-covariance matrices of \mathbf{w} and \mathbf{z} , respectively. $\boldsymbol{\Sigma}_{zw}$ is the covariance between \mathbf{z} and \mathbf{w} . $\boldsymbol{\mu}_w$ and $\boldsymbol{\mu}_z$ are the mean vector of \mathbf{z} and \mathbf{w} respectively.

Relevant space within a model

A concept for reduction of regression models

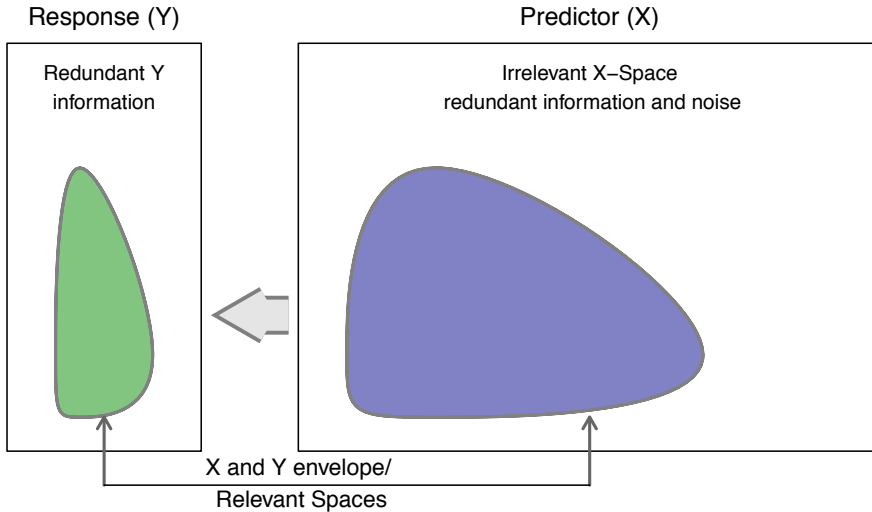


Figure 1: Relevant space in a regression model

Here, the elements of \mathbf{w} and \mathbf{z} are the principal components of responses and predictors, which will respectively be referred to respectively as “response components” and “predictor components”. The column vectors of respective rotation matrices \mathbf{Q} and \mathbf{R} are the eigenvectors corresponding to these principal components. We can write a linear model based on (3) as,

$$\mathbf{w} = \boldsymbol{\mu}_w + \boldsymbol{\alpha}^t(\mathbf{z} - \boldsymbol{\mu}_z) + \boldsymbol{\tau} \quad (4)$$

where, $\boldsymbol{\alpha}^t_{m \times p}$ is a matrix of regression coefficients and $\boldsymbol{\tau}$ is an error term such that $\boldsymbol{\tau} \sim \mathcal{N}(0, \Sigma_{w|z})$.

Following the concept of relevant space, a subset of predictor components can be imagined to span the predictor space. These components can be regarded as relevant predictor components. Næs and Martens [21] introduced the concept of relevant components which was explored further by Helland [10], Næs and Helland [20], Helland and Almøy [12] and Helland [11]. The corresponding eigenvectors were referred to as relevant eigenvectors. A similar logic is introduced by Cook et al. [6] and later by Cook et al. [4] as an envelope which is the space spanned by the relevant eigenvectors [3, pp. 101].

In addition, various simulation studies have been performed with the model based on the concept of relevant subspace. A simulation study by Almøy [1] has used a single response simulation model based on reduced regression and has compared some contemporary multivariate estimators. In recent years Helland et al. [14], Sæbø et al. [25], Helland et al. [13] and Rimal et al. [24] implemented similar simulation examples similar to those we are discussing in this study. This paper, however, presents an elaborate comparison of the prediction using multi-response simulated linear model data. The properties of the simulated data are

varied through different levels of simulation-parameters based on an experimental design. Rimal et al. [24] provide a detailed discussion of the simulation model that we have adopted here. The following section presents the estimators being compared in more detail.

3. Prediction Methods

Partial least squares regression (PLS) and Principal component regression (PCR) have been used in many disciplines such as chemometrics, econometrics, bioinformatics and machine learning, where wide predictor matrices, i.e. p (number of predictors) $> n$ (number of observation) are common. These methods are popular in multivariate analysis, especially for exploratory studies and predictions. In recent years, a concept of envelope introduced by Cook et al. [5] based on the reduction in the regression model was implemented for the development of different estimators. This study compares these prediction methods based on their prediction performance on data simulated with different controlled properties.

Principal Components Regression (PCR): Principal components are the linear combinations of predictor variables such that the transformation makes the new variables uncorrelated. In addition, the variation of the original dataset captured by the new variables is sorted in descending order. In other words, each successive component captures maximum variation left by the preceding components in predictor variables [17]. Principal components regression uses these principal components as a new predictor to explain the variation in the response.

Partial Least Squares (PLS): Two variants of PLS: PLS1 and PLS2 are used for comparison. The first one considers individual response variables separately, i.e. each response is predicted with a single response model, while the latter considers all response variables together. In PLS regression, the components are determined so as to maximize a covariance between response and predictors [9]. R-package `pls` [19] is used for both PCR and PLS methods.

Envelopes: The envelope, introduced by Cook et al. [5], was first used to define response envelope [6] as the smallest subspace in the response space so that the span of regression coefficients lies in that space. Since a multivariate linear regression model contains relevant (material) and irrelevant (immaterial) variation in both response and predictor, the relevant part provides information, while the irrelevant part increases the estimative variation. The concept of the envelope uses the relevant part for estimation while excluding the irrelevant part consequently increasing the efficiency of the model [8].

The concept was later extended to the predictor space, where the predictor envelope was defined [4]. Further Cook and Zhang [7] used envelopes for joint reduction of the responses and predictors and argued that this produced efficiency gains that were greater than those derived by using individual envelopes for either the responses or the predictors separately. All the variants of envelope estimations are based on maximum likelihood estimation. Here we have used predictor envelope (`Xenv`) and simultaneous envelope (`Senv`) for the comparison. R-package `Renvlp` [18] is used for both `Xenv` and `Senv` methods.

3.1. Modification in envelope estimation

Since envelope estimators (Xenv and Senv) are based on maximum likelihood estimation (MLE), it fails to estimate in the case of wide matrices, i.e. $p > n$. To incorporate these methods in our comparison, we have used the principal components (\mathbf{z}) of the predictor variables (\mathbf{x}) as predictors, using the required number of components for capturing 97.5% of the variation in \mathbf{x} for the designs where $p > n$. The new set of variables \mathbf{z} were used for envelope estimation. The regression coefficients ($\hat{\alpha}$) corresponding to these new variables \mathbf{z} were transformed back to obtain coefficients for each predictor variable as,

$$\hat{\beta} = \mathbf{e}_k \hat{\alpha}_k$$

where \mathbf{e}_k is a matrix of eigenvectors with the first k number of components. Only simultaneous envelope allows to specify the dimension of response envelope and all the simulation is based on a single latent dimension of response, so it is fixed at two in the simulation study. In the case of Senv, when the envelope dimension for response is the same as the number of responses, it degenerates to the Xenv method and if the envelope dimension for the predictor is the same as the number of predictors, it degenerates to the standard multivariate linear regression [18].

4. Experimental Design

This study compares prediction methods based on their prediction ability. Data with specific properties are simulated, some of which are easier to predict than others. These data are simulated using the R-package `simrel`, which is discussed in Sæbø et al. [25] and Rimal et al. [24]. Here we have used four different factors to vary the property of the data: a) Number of predictors (p), b) Multicollinearity in predictor variables (γ), c) Correlation in response variables (η) and d) position of predictor components relevant for the response ($relpos$). Using two levels of p , γ and $relpos$ and four levels of η , 32 set of distinct properties are designed for the simulation.

Number of predictors: To observe the performance of the methods on tall and wide predictor matrices, 20 and 250 predictor variables are simulated with the number of observations fixed at 100. Parameter p controls these properties in the `simrel` function.

Multicollinearity in predictor variables: Highly collinear predictors can be explained completely by a few components. The parameter γ in `simrel` controls decline in the eigenvalues of the predictor variables as (5).

$$\lambda_i = e^{-\gamma(i-1)}, \gamma > 0 \text{ and } i = 1, 2, \dots, p \quad (5)$$

Here, $\lambda_i, i = 1, 2, \dots, p$ are eigenvalues of the predictor variables. We have used 0.2 and 0.9 as different levels of γ . The higher the value of γ , the higher the multicollinearity will be, and vice versa.

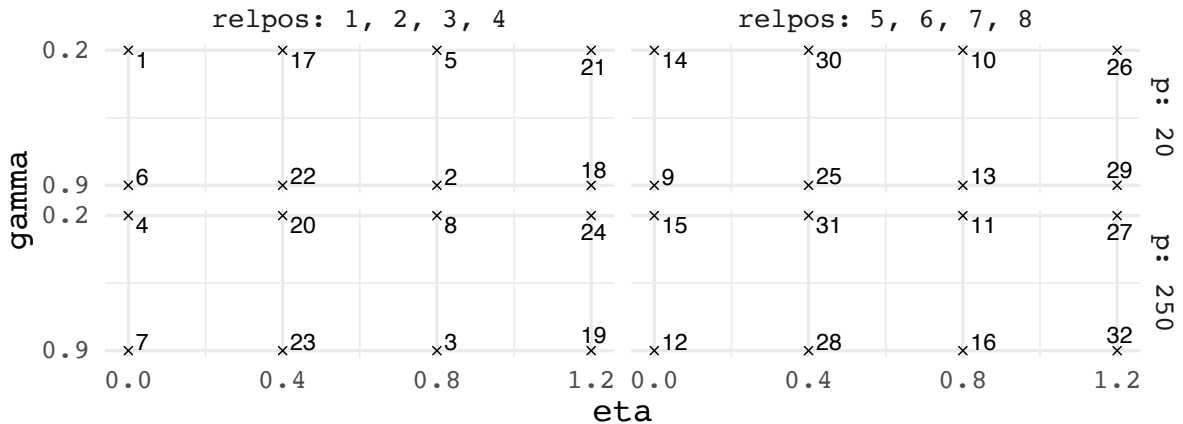


Figure 2: Experimental Design of simulation parameters. Each point represents a unique data property.

Correlation in response variables: Correlation among response variables has been explored to a lesser extent. Here we have tried to explore that part with four levels of correlation in the response variables. We have used the `eta` (η) parameter of `simrel` for controlling the decline in eigenvalues corresponding to the response variables as (6).

$$\kappa_j = e^{-\eta(j-1)}, \eta > 0 \text{ and } j = 1, 2, \dots, m \quad (6)$$

Here, $\kappa_j, i = 1, 2, \dots, m$ are the eigenvalues of the response variables and m is the number of response variables. We have used 0, 0.4, 0.8 and 1.2 as different levels of `eta`. The larger the value of `eta`, the larger will be the correlation will be between response variables and vice versa.

Position of predictor components relevant to the response: The principal components of the predictors are ordered. The first principal component captures most of the variation in the predictors. The second captures most of the remainder left by the first principal component and so on. In highly collinear predictors, the variation captured by the first few components is relatively high. However, if those components are not relevant for the response, prediction becomes difficult [12]. Here, two levels of the positions of these relevant components are used as 1, 2, 3, 4 and 5, 6, 7, 8.

Moreover, a complete factorial design from the levels of the above parameters gave us 32 designs. Each design is associated with a dataset having unique properties. Figure~2, shows all the designs. For each design and prediction method, 50 datasets were simulated as replicates. In total, there were $5 \times 32 \times 50$, i.e. 8000 simulated datasets.

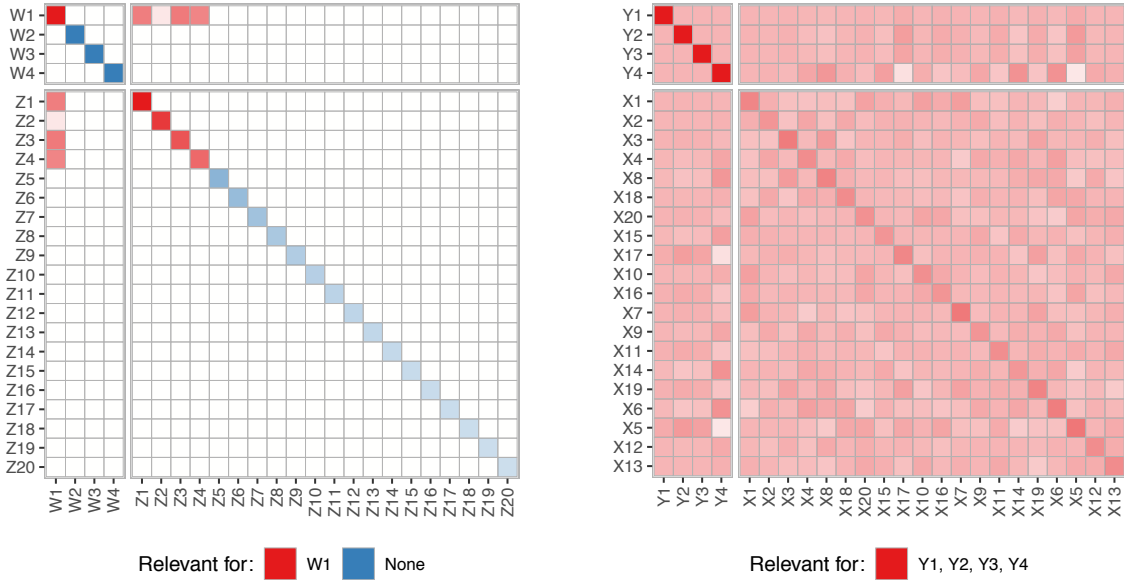


Figure 3: (left) Covariance structure of latent components (right) Covariance structure of predictor and response

Common parameters: Each dataset was simulated with $n = 100$ number of observation and $m = 4$ response variables. Furthermore, the coefficient of determination corresponding to each response components in all the designs is set to 0.8. In addition, we have assumed that there is only one informative response component. Hence, the informative response component is rotated orthogonally together with three uninformative response components to generate four response variables. This spreads out the information in all simulated response variables. For further details on the simulation tool, see [24].

An example of simulation parameters for the first design is as follows:

```
simrel(
  n      = 100,          ## Training samples
  p      = 20,           ## Predictors
  m      = 4,            ## Responses
  q      = 20,           ## Relevant predictors
  relpos = list(c(1, 2, 3, 4)), ## Relevant predictor components index
  eta    = 0,            ## Decay factor of response eigenvalues
  gamma  = 0.2,          ## Decay factor of predictor eigenvalues
  R2     = 0.8,           ## Coefficient of determination
  ypos   = list(c(1, 2, 3, 4)),
  type   = "multivariate"
)
```

The covariance structure of the data simulated with this design in the Figure 3 shows that the predictor

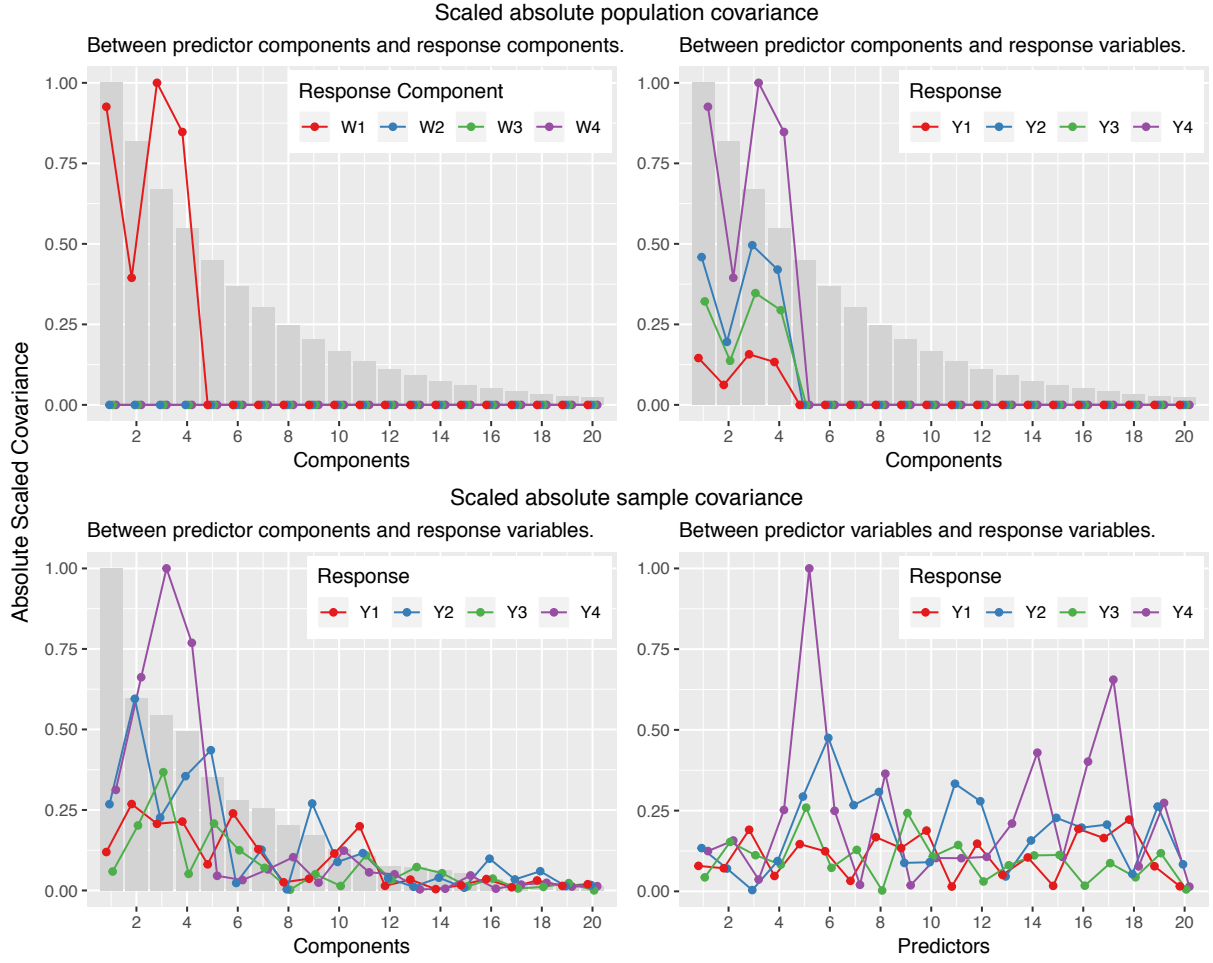


Figure 4: Expected Scaled absolute covariance between predictor components and response components (top left). Expected Scaled absolute covariance between predictor components and response variables (top right). Sample scaled absolute covariance between predictor components and response variables (bottom left). Sample scaled absolute covariance between predictor variables and response variables (bottom right). The bar graph in the background represents eigenvalues corresponding to each component in the population (top plots) and in the sample (bottom plots). One can compare the top-right plot (true covariance of the population) with bottom-left (covariance in the simulated data) which shows a similar pattern for different components.

components at positions 1, 2, 3 and 4 are relevant for the first response component. After the rotation with an orthogonal rotation matrix, all predictor variables are somewhat relevant for all response variables, satisfying other desired properties such as multicollinearity and coefficient of determination. For the same design, Figure 4 (top left) shows that the predictor components 1, 2, 3 and 4 are relevant for the first response component. All other predictor components are irrelevant and all other response components are uninformative. However, due to orthogonal rotation of the informative response component together with uninformative response components, all response variables in the population have similar covariance with the relevant predictor components (Figure 4 (top right)). The sample covariances between the predictor components and predictor variables with response variables are shown in Figure 4 (bottom left) and (bottom right) respectively.

A similar description can be made for all 32 designs, where each of the designs holds the properties of the data they simulate. These data are used by the prediction methods discussed in the previous section. Each prediction method is given independently simulated datasets in order to give them an equal opportunity to capture the dynamics in the data.

5. Basis of comparison

This study focuses mainly on the prediction performance of the methods with an emphasis specifically on the interaction between the properties of the data controlled by the simulation parameters and the prediction methods. The prediction performance is measured based on the following:

- a) The average prediction error that a method can give using an arbitrary number of components and
- b) The average number of components used by the method to give the minimum prediction error

Let us define,

$$\mathcal{P}\mathcal{E}_{ijkl} = \frac{1}{\sigma_{y_{ij}|x}^2} E \left[\left(\beta_{ij} - \hat{\beta}_{ijkl} \right)^t (\Sigma_{xx})_i \left(\beta_{ij} - \hat{\beta}_{ijkl} \right) \right] + 1 \quad (7)$$

as a prediction error of response $j = 1, \dots, 4$ for a given design $i = 1, 2, \dots, 32$ and method $k = 1(\text{PCR}), \dots, 5(\text{Senv})$ using $l = 0, \dots, 10$ number of components. Here, $(\Sigma_{xx})_i$ is the true covariance matrix of the predictors, unique for a particular design i and $\sigma_{y_{ij}|x}^2$ for response $j = 1, \dots, m$ is the true model error. Here prediction error is scaled by the true model error to remove the effects of influencing residual variances. Since both the expectation and the variance of $\hat{\beta}$ are unknown, the prediction error is estimated using data from 50 replications as follows,

$$\widehat{\mathcal{P}\mathcal{E}}_{ijkl} = \frac{1}{\sigma_{y_{ij}|x}^2} \sum_{r=0}^{50} \left[\left(\beta_{ij} - \hat{\beta}_{ijklr} \right)^t (\Sigma_{xx})_i \left(\beta_{ij} - \hat{\beta}_{ijklr} \right) \right] + 1 \quad (8)$$

where $\widehat{\mathcal{P}\mathcal{E}}_{ijkl}$ is the estimated prediction error averaged over $r = 50$ replicates.

The following section focuses on the data for the estimation of these prediction errors that are used for the two models discussed above in a) and b) of this section.

6. Data Preparation

A dataset for estimating (7) is obtained from simulation which contains a) five factors corresponding to simulation parameters, b) prediction methods, c) number of components, d) replications and e) prediction error for four responses. The prediction error is computed using predictor components ranging from 0 to 10 for every 50 replicates as,

$$(\widehat{\mathcal{P}\mathcal{E}}_o)_{ijklr} = \frac{1}{\sigma_{y_{ij}|x}^2} \left[\left(\beta_{ij} - \hat{\beta}_{ijklr} \right)^t (\Sigma_{xx})_i \left(\beta_{ij} - \hat{\beta}_{ijklr} \right) \right] + 1$$

Thus there are 32 (designs) \times 5 (methods) \times 11 (number of components) \times 50 (replications), i.e. 88000 observations corresponding to the response variables from Y1 to Y4.

Since our discussions focus on the average minimum prediction error that a method can obtain and the average number of components they use to get the minimum prediction error in each replicates, the dataset discussed above is summarized as constructing the following two smaller datasets. Let us call them *Error Dataset* and *Component Dataset*.

Error Dataset: For each prediction method, design and response, an average prediction error is computed over all replicates for each component. Next, a component that gives the minimum of this average prediction error is selected, i.e.,

$$l_o = \operatorname{argmin}_l \left[\frac{1}{50} \sum_{i=1}^{50} (\mathcal{PE}_o)_{ijklr} \right] \quad (9)$$

Using the component l_o , a dataset of $(\mathcal{PE}_o)_{ijklr}$ is used as the *Error Dataset*. Let $\mathbf{u}_{(8000 \times 4)} = (u_j)$ for $j = 1, \dots, 4$ be the outcome variables measuring the prediction error corresponding to the response number j in the context of this dataset.

Component Dataset: The number of components that gives the minimum prediction error in each replication is referred to as the *Component Dataset*, i.e.,

$$l_o = \operatorname{argmin}_l [\mathcal{PE}_{ijklr}] \quad (10)$$

Here l_o is the number of components that gives minimum prediction error $(\mathcal{PE}_o)_{ijklr}$ for design i , response j , method k and replicate r . Let $\mathbf{v}_{(8000 \times 4)} = (v_j)$ for $j = 1, \dots, 4$ be the outcome variables measuring the number of components used for minimum prediction error corresponding to the response j in the context of this dataset.

7. Exploration

This section explores the variation in the *error dataset* and the *component dataset* for which we have used Principal Component Analysis (PCA). Let \mathbf{t}_u and \mathbf{t}_v be the principal component score sets corresponding to PCA run on the \mathbf{u} and \mathbf{v} matrices respectively. The scores density in Figure-5 corresponds to the first principal component of \mathbf{u} , i.e. the first column of \mathbf{t}_u .

Since higher prediction errors correspond to high scores, the plot shows that the PCR, PLS1 and PLS2 methods are influenced by the two levels of the position of relevant predictor components. When the relevant predictors are at positions 5, 6, 7, 8, the eigenvalues corresponding to them are relatively smaller. This also suggests that PCR, PLS1 and PLS2 depend greatly on the position of the relevant components, and

the variation of these components affects their prediction performance. However, the envelope methods appeared to be less influenced by `relops` in this regard.



Figure 5: Scores density corresponding to first principal component of *error dataset* (**u**) subdivided by `methods`, `gamma` and `eta` and grouped by `relops`.

In addition, the plot also shows that the effect of `gamma`, i.e., the level of multicollinearity, has a lesser effect when the relevant predictors are at positions 1, 2, 3, 4. This indicates that the methods are somewhat robust for handling collinear predictors. Nevertheless, when the relevant predictors are at positions 5, 6, 7, 8, high multicollinearity results in a small variance of these relevant components and consequently yields poor prediction. This is in accordance with the findings of Helland and Almøy [12].

Furthermore, the density curves for PCR, PLS1 and PLS2 are similar for different levels of `eta`, i.e., the factor controlling the correlation between responses. However, the envelope models have been shown to have distinct interactions between the positions of relevant components (`relops`) and `eta`. Here higher levels of `eta` have yielded higher scores and clear separation between two levels of `relops`.

In the case of high multicollinearity, envelope methods have resulted in some large outliers indicating that in some cases that the methods can result in giving an unexpected prediction.

In Figure 6, the higher scores suggest that methods have used a larger number of components to give minimum prediction error. The plot also shows that the relevant predictor components at 5, 6, 7, 8 give larger

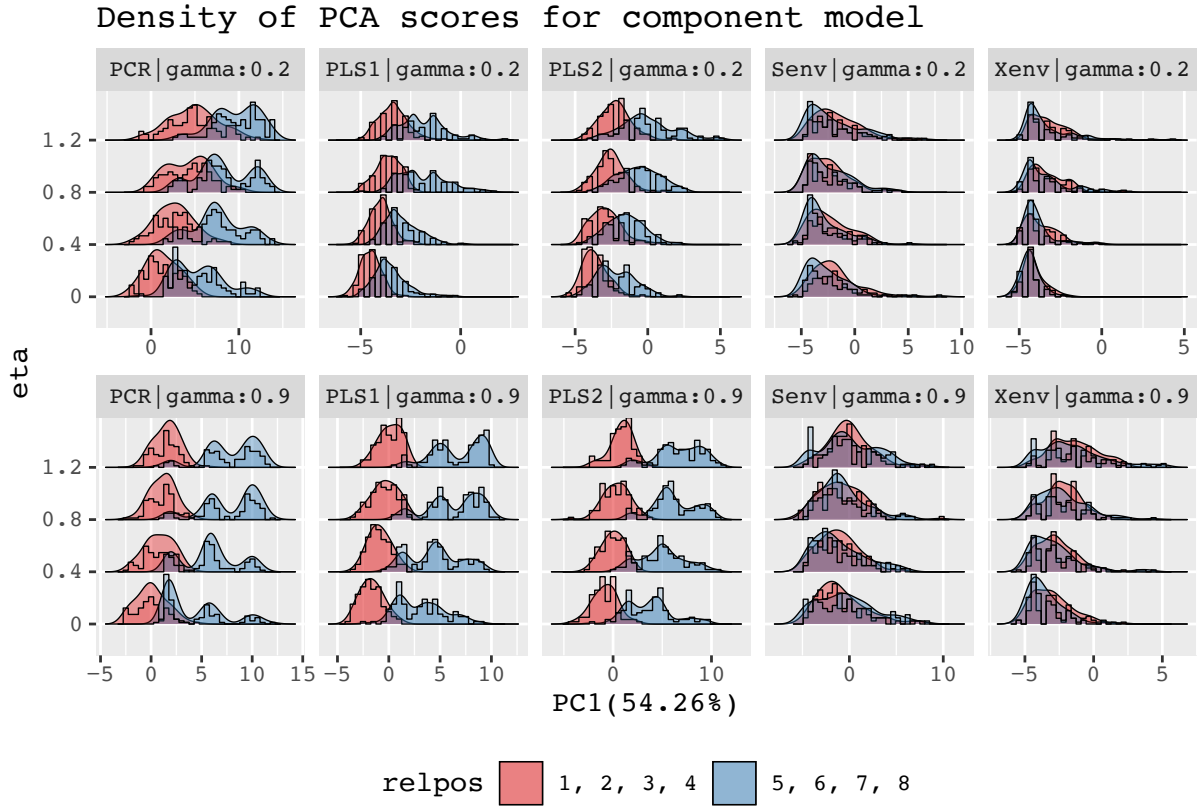


Figure 6: Score density corresponding to first principal component of *component dataset* (**v**) subdivided by **methods**, **gamma** and **eta** and grouped by **relpos**.

prediction errors than those in positions 1, 2, 3, 4. The pattern is more distinct in large multicollinearity cases and PCR and PLS methods. Both the envelope methods have shown equally enhanced performance at both levels of **relpos** and **gamma**. However, for data with low multicollinearity ($\gamma = 0.2$), the envelope methods have used a lesser number of components on average than in the high multicollinearity cases to achieve minimum prediction error.

8. Statistical Analysis

This section has modelled the *error data* and the *component data* as a function of the simulation parameters to better understand the connection between data properties and prediction methods using multivariate analysis of variation (MANOVA).

Let us consider a model with third order interaction of the simulation parameters (**p**, **gamma**, **eta** and **relpos**) and Methods as in (11) and (12) using datasets **u** and **v**, respectively. Let us refer them as the *error model* and the *component model*.

Error Model:

$$\mathbf{u}_{abcdef} = \boldsymbol{\mu}_u + (\mathbf{p}_a + \mathbf{gamma}_b + \mathbf{eta}_c + \mathbf{relpos}_d + \mathbf{Methods}_e)^3 + (\boldsymbol{\varepsilon}_u)_{abcdef} \quad (11)$$

Component Model:

$$\mathbf{v}_{abcdef} = \boldsymbol{\mu}_v + (\text{p}_a + \text{gamma}_b + \text{eta}_c + \text{re1pos}_d + \text{Methods}_e)^3 + (\boldsymbol{\varepsilon}_v)_{abcdef} \quad (12)$$

where, \mathbf{u}_{abcdef} is a vector of prediction errors in the *error model* and \mathbf{v}_{abcdef} is a vector of the number of components used by a method to obtain minimum prediction error in the *component model*.

Although there are several test-statistics for MANOVA, all are essentially equivalent for large samples [16]. Here we will use Pillai's trace statistic which is defined as,

$$\text{Pillai statistic} = \text{tr} \left[(\mathbf{E} + \mathbf{H})^{-1} \mathbf{H} \right] = \sum_{i=1}^m \frac{\nu_i}{1 + \nu_i} \quad (13)$$

Here the matrix \mathbf{H} holds between-sum-of-squares and sum-of-products for each of the predictors. The matrix \mathbf{E} has a within the sum of squares and sum of products for each of the predictors. ν_i represents the eigenvalues corresponding to $\mathbf{E}^{-1} \mathbf{H}$ [23].

For both the models (11) and (12), Pillai's trace statistic is used for accessing the effect of each factor and returns an F-value for the strength of their significance. Figure 7 plots the Pillai's trace statistics as bars with corresponding F-values as text labels for both models.

Error Model: Figure 7 (left) shows the Pillai's trace statistic for factors of the *error model*. The main effect of Method followed by re1pos, eta and gamma have largest influence on the model. A highly significant two-factor interaction of Method with eta followed by re1pos and gamma clearly shows that methods perform differently for different levels of these data properties. The significant third order interaction between Method, eta and gamma suggests that the performance of a method differs for a given level of multicollinearity and the correlation between the responses. Since only some methods consider modelling predictor and response together, the prediction is affected by the level of correlation between the responses (eta) for a given method.

Component Model: Figure 7 (right) shows the Pillai's trace statistic for factors of the *component model*. As in the *error model*, the main effects of the Method, re1pos, gamma and eta have a significantly large effect on the number of components that a method has used to obtain minimum prediction error. The two-factor interactions of Method with simulation parameters are larger in this case. This shows that the Methods and these interactions have a larger effect on the use of the number of component than the prediction error itself. In addition, a similar significant high third-order interaction as found in the *error model* is also observed in this model.

The following section will continue to explore the effects of different levels of the factors in the case of these interactions.

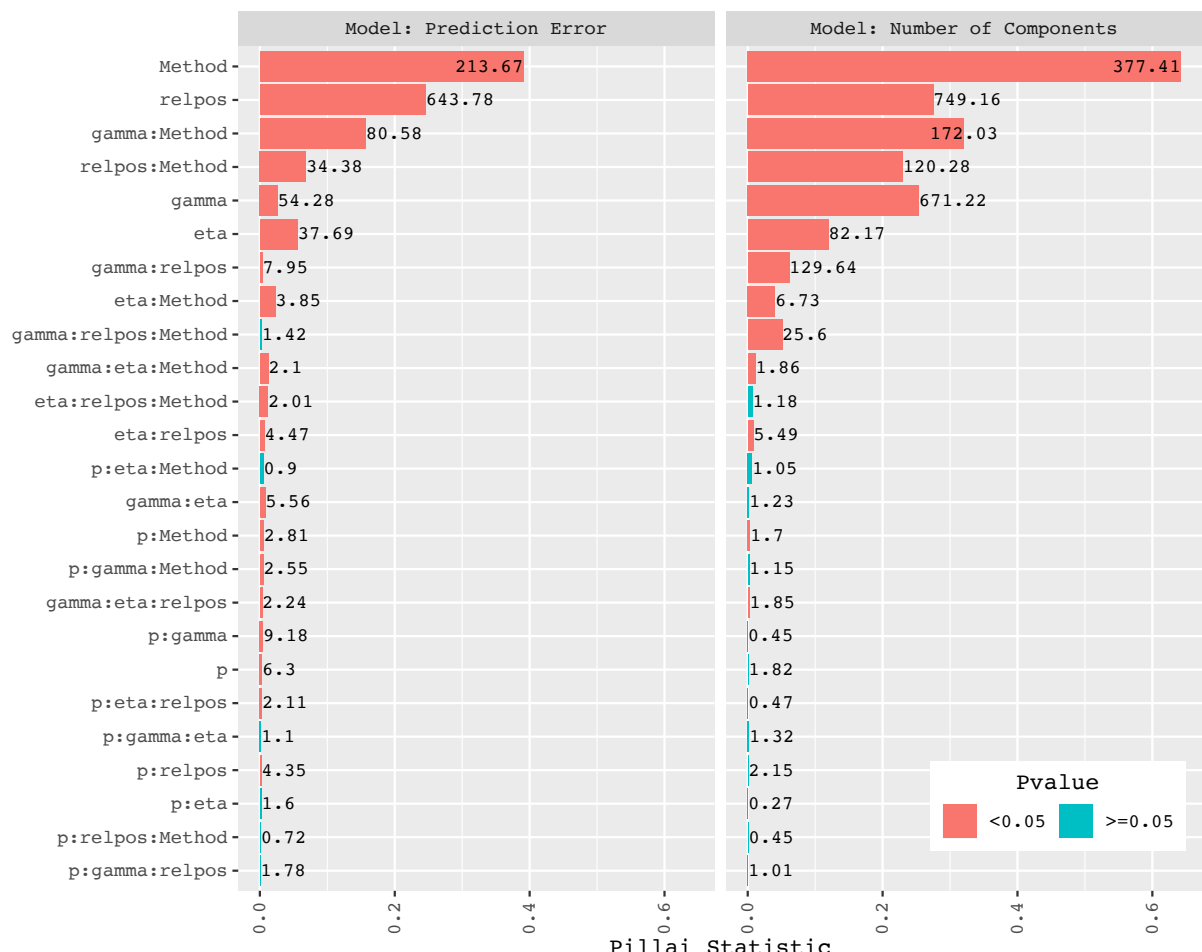


Figure 7: Pillai Statistic and F-value for the MANOVA model. The bar represents the Pillai Statistic and the text labels are F-value for the corresponding factor.

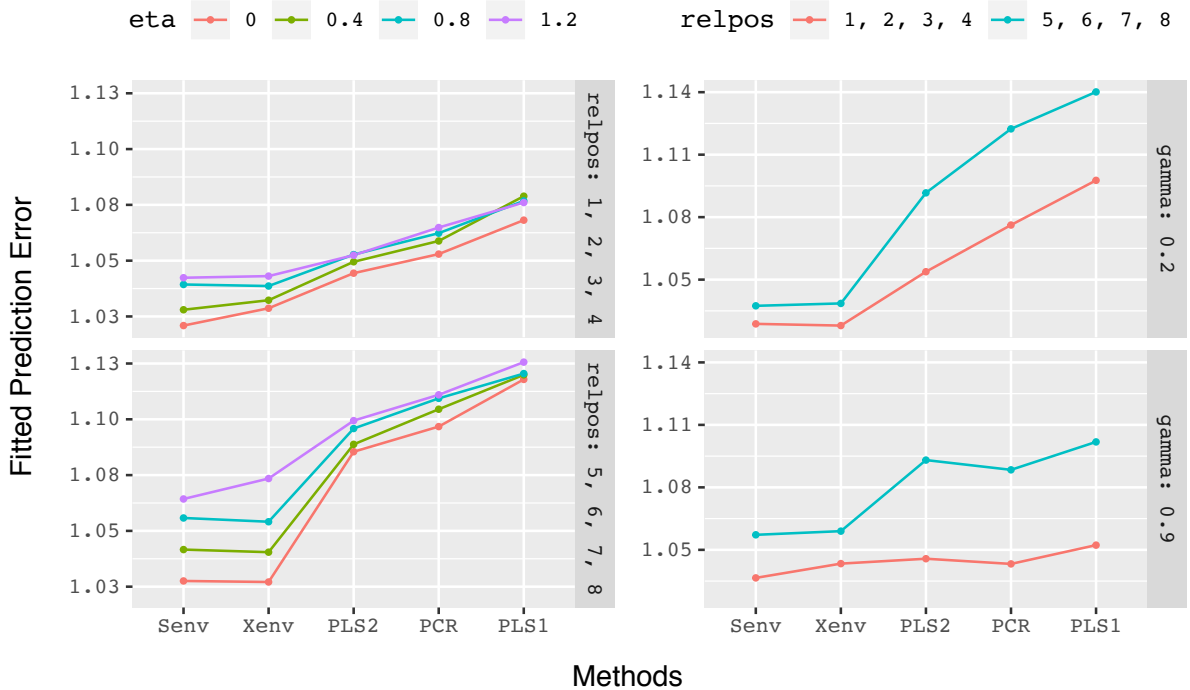


Figure 8: Effect plot of some interactions of the multivariate linear model of prediction error

8.1. Effect Analysis of Error Model

The large difference in the prediction error for the envelope models in Figure 8 (left) is intensified when the position of the relevant predictor is at 5, 6, 7, 8. The results also show that the envelope methods are more sensitive to the levels of eta than the rest of the methods. In the case of PCR and PLS, the difference in the effect of levels of eta is small.

In Figure 8 (right), we can see that the multicollinearity (controlled by gamma) has affected all the methods. However, envelope methods have better performance on low multicollinearity, as opposed to high multicollinearity, and PCR, PLS1 and PLS2 are robust for high multicollinearity. Despite handling high multicollinearity, these methods have higher prediction error in both cases of multicollinearity than the envelope methods.

8.2. Effect Analysis of Component Model

Unlike for prediction errors, Figure 9 (left) shows that the number of components used by the methods to obtain minimum prediction error is less affected by the levels of eta. All methods appear to use on average more components when eta increases. Envelope methods are able to obtain minimum prediction error by using components ranging from 1 to 3 in both the cases of relpos. This value is much higher in the case of PCR as its prediction is based only on the principal components of the predictor matrix. The number of components used by this method ranges from 3 to 5 when relevant components are at positions 1, 2, 3, 4 and 5 to 8 when relevant components are at positions 5, 6, 7, 8.

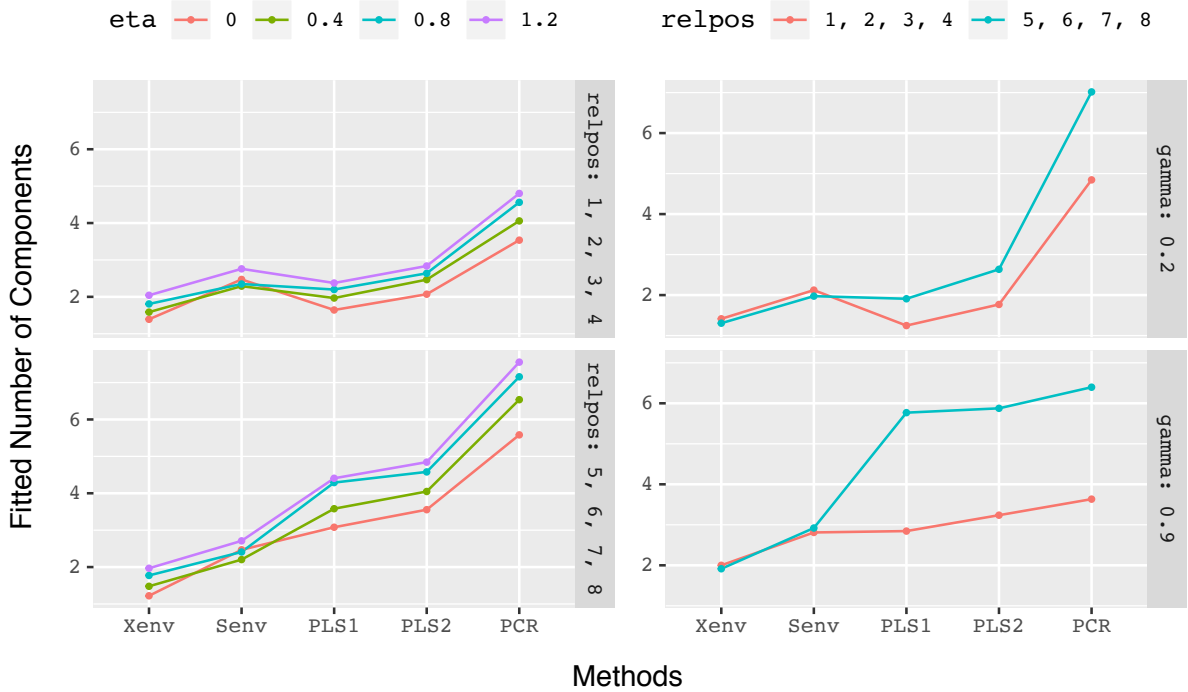


Figure 9: Effect plot of some interactions of the multivariate linear model of the number of components to get minimum prediction error

When relevant components are at position 5, 6, 7, 8, the eigenvalues of relevant predictors becomes smaller and responses are relatively difficult to predict. This becomes more critical for high multicollinearity cases. Figure 9 (right) shows that the envelope methods are less influenced by the level of `relpos` and are particularly better in achieving minimum prediction error using a fewer number of components than other methods.

9. Examples

In addition to the analysis with the simulated data, the following two examples explore the prediction performance of the methods using real datasets. Since both examples have wide predictor matrices, principal components explaining 97.5% of the variation in them are used for envelope methods. The coefficients were transformed back after the estimation.

9.1. Raman spectra analysis of contents of polyunsaturated fatty acids (PUFA)

This dataset contains 44 training samples and 25 test samples of fatty acid information expressed as: a) percentage of total sample weight and b) percentage of total fat content. The dataset is borrowed from Næs et al. [22] where more information can be found. The samples were analysed using Raman spectroscopy from which 1096 wavelength variables were obtained as predictors. Raman spectroscopy provides detailed chemical information from minor components in food. The aim of this example is to compare how well the

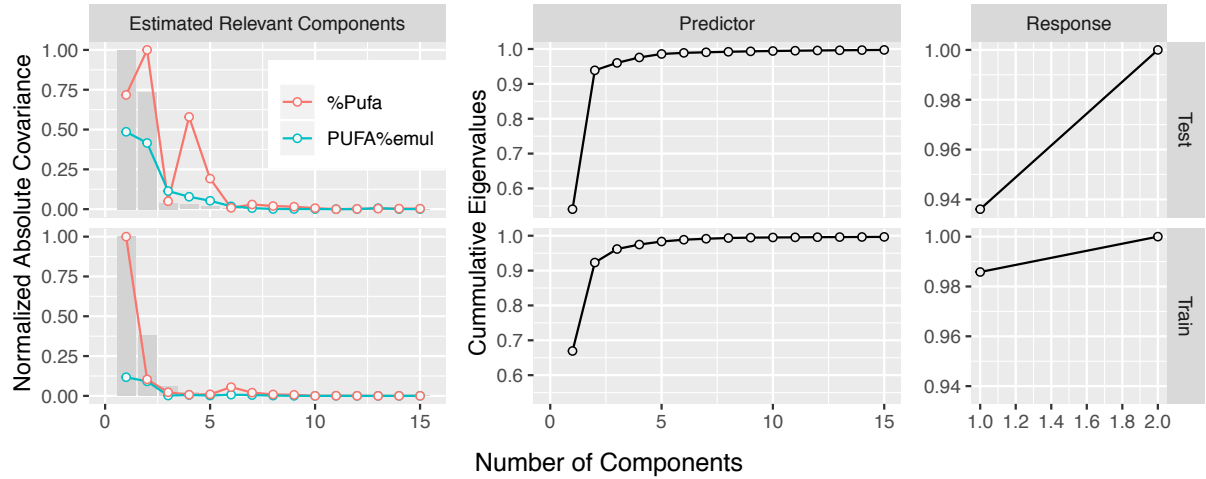


Figure 10: (Left) Bar represents the eigenvalues corresponding to Raman Spectra. The points and line are the covariances between response and the principal components of Raman Spectra. All the values are normalized to scale from 0 to 1. (Middle) Cumulative sum of eigenvalues corresponding to predictors. (Right) Cumulative sum of eigenvalues corresponding to responses. The top and bottom row corresponds to test and training datasets respectively.

prediction methods that we have considered are able to predict the contents of PUFA using these Raman spectra.

Figure 10 (left) shows that the first few predictor components are somewhat correlated with response variables. In addition, the most variation in predictors is explained by less than five components (middle). Further, the response variables are highly correlated, suggesting that a single latent dimension explains most of the variation (right). We may therefore also believe that the relevant latent space in the response matrix is of dimension one. This resembles the Design 19 (Figure 2) from our simulation.

Using a range of components from 1 to 15, regression models were fitted using each of the methods. The fitted models were used to predict the test observation, and the root mean squared error of prediction (RMSEP) was calculated. Figure 11 shows that PLS2 obtained a minimum prediction error of 3.783 using 9 components in the case of response %Pufa, while PLS1 obtained a minimum prediction error of 1.308 using 11 components in the case of response PUFA%emul. However, the figure also shows that both envelope methods have reached to almost minimum prediction error in fewer number of components. This pattern is also visible in the simulation results (Figure 9).

9.2. Example-2: NIR spectra of biscuit dough

The dataset consists of 700 wavelengths of NIR spectra (1100–2498 nm in steps of 2 nm) that were used as predictor variables. There are four response variables corresponding to the yield percentages of (a) fat, (b) sucrose, (c) flour and (d) water. The measurements were taken from 40 training observation of biscuit dough. A separate set of 32 samples created and measured on different occasions were used as test observations. The dataset is borrowed from Indahl [15] where further information can be obtained.

Figure 12 (left) shows that the first predictor component has the largest variance and also has large covariance

Prediction Error per Response

For each prediction method

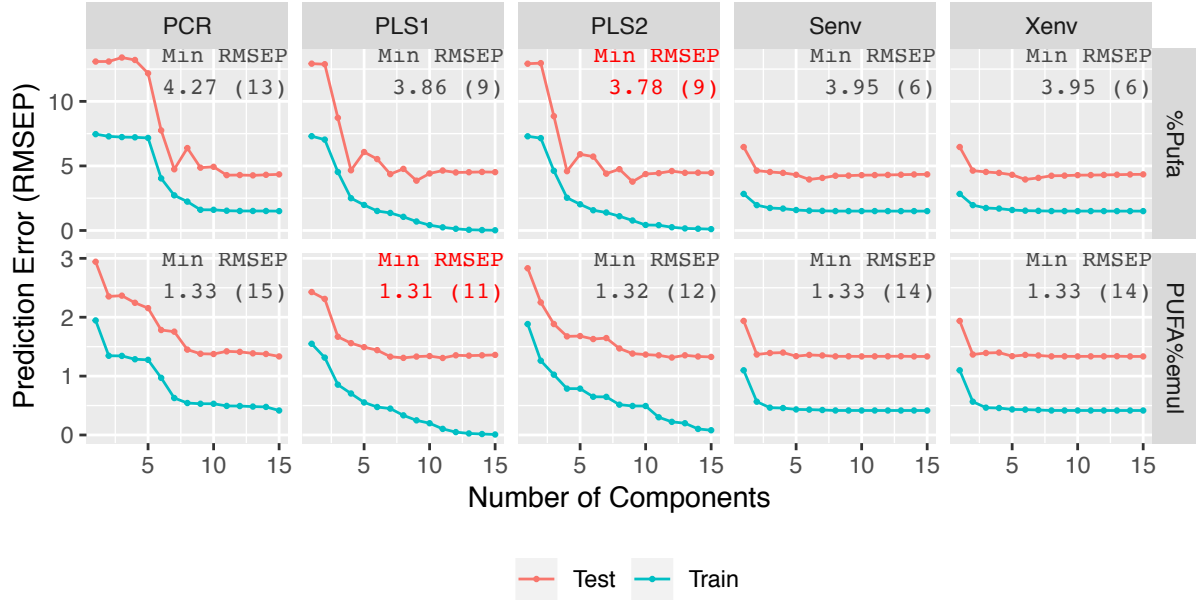


Figure 11: Prediction Error of different prediction methods using different number of components.

with all response variables. The second component, however, has larger variance (middle) than the succeeding components but has a small covariance with all the responses, which indicates that the component is less relevant for any of the responses. In addition, two response components have explained most of the variation in response variables (right). This structure is also somewhat similar to Design 19, although it is uncertain whether the dimension of the relevant space in the response matrix is larger than one.

Figure 13 (corresponding to Figure 11) shows the root mean squared error for both test and train prediction of the biscuit dough data. Here four different methods have minimum test prediction error for the four responses. As the structure of the data is similar to that of the first example, the pattern in the prediction is also similar for all methods.

The prediction performance on the test data of the envelope methods appears to be more stable compared to the PCR and PLS methods. Furthermore, the envelope methods achieve good performance generally using fewer components, which is in accordance with Figure 6.

10. Discussions and Conclusion

Analysis using both simulated data and real data has shown that the envelope methods are more stable, less influenced by `relpos` and `gamma` and in general, performed better than PCR and PLS methods. These methods are also found to be less dependent on the number of components.

Since the facet in the Figures 5 and 6 have their own scales, despite having some large prediction errors seen

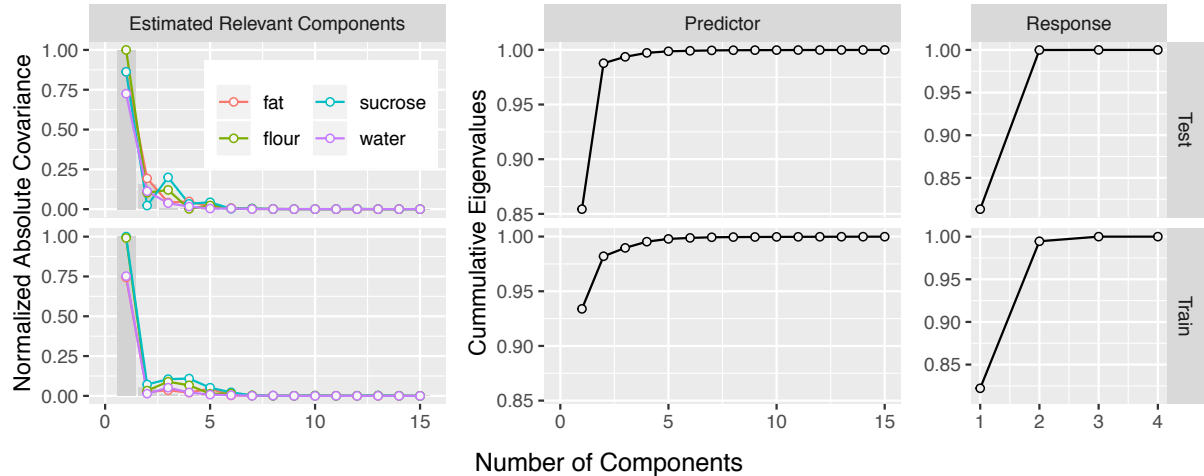


Figure 12: (Left) Bar represents the eigenvalues corresponding to NIR Spectra. The points and line are the covariances between response and the principal components of NIR Spectra. All the values are normalized to scale from 0 to 1. (Middle) Cumulative sum of eigenvalues corresponding to predictors. (Right) Cumulative sum of eigenvalues corresponding to responses.

at the right tail, envelope methods still have a smaller prediction error and have used a fewer number of components than the other methods.

Particularly in the case of the simultaneous envelope, since users can specify the number of dimension for the response envelope, the method can leverage the relevant space of response while PCR, PLS and Xenv are constrained to play only on predictor space. *Since the simulation is based on a single latent component of the response variables, this might have given some advantage for the simultaneous envelope.*

Furthermore, we have fixed the coefficient of determination (R^2) as a constant throughout all the designs. Initial simulations (not shown) indicated that low R^2 affects all methods in a similar manner and that the MANOVA is highly dominated by R^2 . Keeping the value of R^2 fixed has allowed us to analyze other factors properly.

Two clear comments can be made about the effect of correlation of response on the prediction methods. The highly correlated response has shown the highest prediction error in general and the effect is most distinct in envelope methods. Since the envelope methods identify the relevant space as the span of relevant eigenvectors, the methods are able to obtain the minimum average prediction error by using a lesser number of components for all levels of eta.

To our knowledge, the effect of correlation in the response on PCR and PLS methods has been explored only to a limited extent. In this regards, it is interesting to see that these methods have applied a large number of components and returned a larger prediction error than envelope methods in the case of highly correlated responses. To fully understand the effect of eta, it is necessary to study the estimation performance of these methods with different numbers of components.

In addition, since using principal components or actual variables as predictors in envelope methods has shown similar results, we have used principal components that have explained 97.5% of the variation as

Prediction Error per Response

For each prediction method

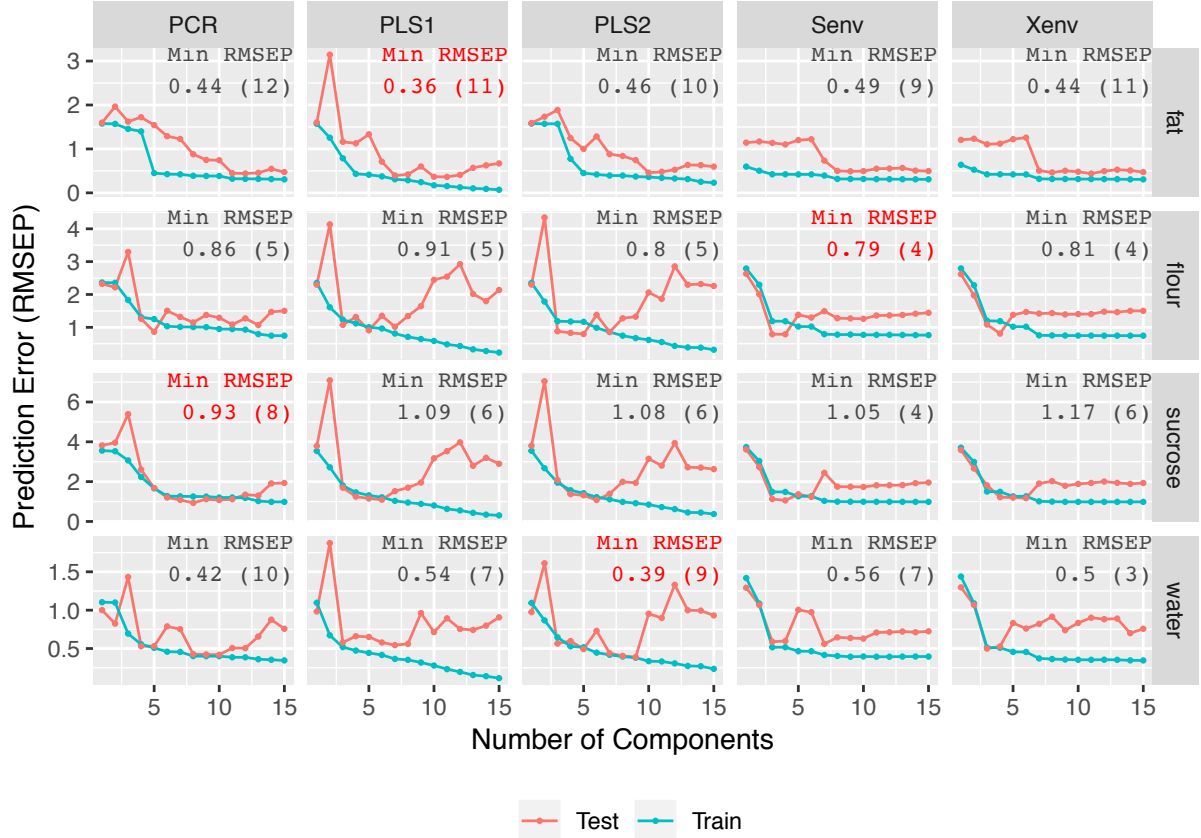


Figure 13: Prediction Error of different prediction methods using different number of components.

mentioned previously in the cases of envelope methods for the designs where $p > n$. As the envelope methods are based on MLE, this can be an alternative way of using the methods in data with wide predictors. The results from this study will help researchers to understand these methods for their performance in various linear model data and encourage them to use newly developed methods such as the envelopes. Since this study has focused entirely on prediction performance, further analysis of the estimative properties of these methods is required. A study of estimation error and the performance of methods on the non-optimal number of components can give a deeper understanding of these methods.

A shiny application [2] is available at <http://therimalaya.shinyapps.io/Comparison> where all the results related to this study can be visualized. In addition, a GitHub repository at <https://github.com/therimalaya/03-prediction-comparison> can be used to reproduce this study.

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