A Two-Step Regression Method with Connections to Partial Least Squares and the Growth Curve Model

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Abstract

Prediction of a continuous response variable from background data is considered. The independent prediction variable data may have a collinear structure and comprise group effects. A new two-step regression method inspired by PLS (partial least squares regression) is proposed. The proposed new method is coupled to a novel application of the Cayley-Hamilton theorem and a two-step estimation procedure. In the two-step approach, the first step summarizes the information in the predictors via a bilinear model. The bilinear model has a Krylov structured within-individuals design matrix, which is closely linked to PLS, and a between-individuals design matrix, which allows the model to handle complex structures, e.g. group effects. The second step is the prediction step, where conditional expectation is used. The close relation between the two-step method and PLS gives new insight into PLS; i.e. PLS can be considered as an algorithm for generating a Krylov structured sequence to approximate the inverse of the covariance matrix of the predictors. Compared with classical PLS, the new two-step method is a non-algorithmic approach. The bilinear model used in the first step gives a greater modelling flexibility than classical PLS. The proposed new two-step method has been extended to handle grouped data, especially data with different mean levels and with nested mean structures. Correspondingly, the new two-step method uses bilinear models with a structure similar to that of the classical growth curve model and the extended growth curve model, but with design matrices which are unknown. Given that the covariance between the predictors and the response is known, the explicit maximum likelihood estimators (MLEs) for the dispersion and mean of the predictors have all been derived. Real silage spectra data have been used to justify and illustrate the two-step method.

Keywords: A Two-Step Regression Method, Growth Curve Model, Krylov Space, MLE, PLS.

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Dedication

To Jianxin and Yichen
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This thesis is based on the work documented in the following papers, referred to by Roman numerals.


IV Ying Li, Peter Udén and Dietrich von Rosen (2014). A two-step method for group data with connections to the extended growth model and PLS. *Submitted.*

The papers have been reprinted in the thesis with the permission of the publishers.
1 Introduction

The prediction problem of a response variable based on some multivariate variables is at the core of statistical applications. One common choice of prediction method is to use the ordinary least squares (OLS) estimator. The Gauss-Markov theory asserts that the least squares estimator is BLUE (the best linear unbiased estimator). However, unbiasedness is not necessarily a wise criterion for estimation, especially when it concerns prediction. The prediction accuracy is related to the mean squares error (MSE) of the estimators and it is the sum of the variance and the squared bias. The MSE of the OLS estimator is the smallest compared with the MSEs of all the other linear unbiased estimators. However, there exist estimators with a small bias but a large variance reduction which have better overall prediction accuracy than that of an OLS estimator. For example, when the variables are “collinear” or “near-collinear”, the prediction accuracy may be poor.

Collinearity refers in a strict sense to the presence of exact linear relationships within a set of variables, typically a set of explanatory (predictor) variables used in a regression-type model. In common statistical language collinearity also allows near-collinearity, when the variables are close to being linearly related, i.e. when their correlation matrix is near-singular, in other words, when the data are ill-conditioned (Sundberg, 2002). Several methods have been proposed for handling the problems connected with collinear data, including ridge regression (RR), the least absolute shrinkage and selection operator (lasso), principal component regression (PCR) and partial least squares regression (PLSR); for a review see Brown (1993) and Sundberg (1999). In particular, PLS, which is the dominant prediction method in chemometrics, is considered in some detail in the thesis. In the literature, some people prefer using the abbreviation PLSR instead of PLS, thereby emphasising that this is a regression method. However, we will use PLS in this thesis.

Partial least squares regression originated from the non-linear iterative partial least squares (NIPALS) algorithm under the concept of “soft modelling” developed by H. Wold (Wold, 1966). “Soft modelling” means building models without any assumptions about the underlying distribution, in contrast to the traditional culture of (“hard”) model building, and is used especially for complex situations where any prior information is scarce. “Soft modelling” is proposed as a means of complementing the weak points of the ordinary statistical modelling culture, where the statistical analysis is started by formulating a parametric model for the data, including a number of essential assumptions. Thereafter, model criticism takes place, addressing whether the data suggest a minor or major modification of the model. One criticism against the “ordinary culture” maintains that data will often point with equal emphasis to several possible models without any specific distribution preference. However, in a defence of “hard modelling”, in practice, using one particular model will not sacrifice much of the information in the data. Furthermore, several statistical methods usually possess certain robustness properties with respect to the certain model assumptions. It is questionable whether “soft modelling”,
with its loose modelling concept, will overcome some of the problems of “hard modelling”. Moreover, by applying the “soft modelling” ideas, one gains little theoretical understanding of the underlying problems. It is very difficult to assess the methods in a general context without a precise modelling concept. On the other hand, employing “soft modelling”, the practitioners apply the comfortable notion that they can collect a batch of data without having to worry too much about how the data have been collected or what past knowledge there is. They can apply some algorithms, e.g. PLS, from soft sciences with the assurance that after some fine tuning they will have a good predictor for future unspecified purposes. The opposing view is that “hard modelling” is of paramount importance (Brown, 1993).

Nevertheless, we cannot deny the fact that practical success has been achieved by applying partial least squares regression in particular within “soft modelling”. Therefore, for mathematical statisticians, instead of debating the concept theoretically, it would be wise to keep one’s mind open, to try to understand the techniques developed “outside” (in the surroundings of “hard modelling”) and to find the harmony that exists between the new and the old, thereby refreshing our theory and improving development. The first attempt to take PLS into the ordinary statistical culture was made by Helland (1988, 1990) who defined the population algorithm by emphasising the distinction between the parameters and observations. Recently, Cook et al. (2013) found a close connection between PLS and the envelope model, which is a new general methodology for dimension reduction. There are also a number of other important works which try to clarify PLS theoretically, for example those by Stone and Brooks (1990), Frank and Friedman (1993), and Butler and Denham (2000).

The overall goal of the present thesis is to cast PLS in a probability model framework. Then, under such a framework, we aim to develop new methods for general cases. In particular, we show the connections between the population version of PLS and a new proposed two-step regression method. The two-step method is based on strict stochastic assumptions. In the first step, information in the explanatory variable is extracted with the help of a multivariate linear model. In the second step, the prediction step, a conditional approach is applied. In a special setting, the two-step method produces the same predictor as PLS.

The aims of the thesis can be summarized as follows:

1. to develop a new PLS-inspired method which combines efficient prediction in collinear cases with a well-defined statistical model;

2. to extend the new developed method to more complex data structures which comprise group effects, which in turn may be nested.

We begin in Section 2 by presenting the background of the thesis, including an introduction to a few basic models and a review of shrinkage methods, PLS and growth curve models. The papers which this thesis is based on, are
summarized in Section 3. The research contributions made by this thesis and proposals for future research are discussed in the last section.
2 Background

2.1 Basic model assumptions

Let \((x', y')'\) be a \((p + 1)\)-dimensional random vector, with a joint multivariate distribution with \(E[x] = \mu_x\) and \(E[y] = \mu_y\), where \(E[\cdot]\) denotes the mean, \(D[x] = \Sigma\) (supposed to be positive definite), where \(D[\cdot]\) denotes the dispersion (variance), and \(C[x, y] = \omega\), where \(C[\cdot]\) denotes the covariance, and under the usual normality assumption we may write

\[
\begin{pmatrix} x \\ y \end{pmatrix} \sim N_{(p+1)} \left( \begin{pmatrix} \mu_x \\ \mu_y \end{pmatrix}, \begin{pmatrix} \Sigma & \omega \\ \omega' & \sigma^2_y \end{pmatrix} \right). \tag{2.1}
\]

The purpose is to predict \(y\) from \(x\) on the basis of new observations. When all the parameters are known, even without the assumption of normality, the best linear predictor is the conditional expectation of \(y\) given \(x\), i.e.

\[
\hat{y} = E[y|x] = \beta'(x - \mu_x) + \mu_y, \quad \beta = \Sigma^{-1}\omega, \tag{2.2}
\]

where \(E[y|x]\) is the conditional expectation.

2.2 PLS

Partial least squares was first presented as the non-linear iterative partial least squares (NIPALS) algorithm under the concept of “soft modelling”, which was developed by H. Wold (Wold, 1966) and applied to a block of variables in order to find principal components. Since then, H. Wold has presented partial least squares path modelling for the analysis of several sets of variables linked in a path diagram. Today the development of partial least squares may be considered as heading in two main directions. In one direction, the soft modelling concept continues to be used as an alternative algorithm to structural equation models and is finding applications in social sciences. In the other direction partial least squares is used as a new regression method, i.e. partial least squares regression (PLS), and is often applied in chemometrics. In this thesis, we will focus on PLS as a regression method.

2.2.1 The sample version of PLS

Partial least squares regression was mainly developed by S. Wold and H. Martens at the end of the 1970s as a prediction method within the chemometric field. Since then it has been heavily promoted and there are many application papers in the literature, despite the fact that it is only algorithmically defined. The original algorithm of PLS is formulated as follows (Wold et al., 1983).

1. Start with data \((X, y)\) of dimension \((p + 1) \times n\),

\[
E_0 = X - \bar{X} \quad f_0 = y - \bar{y}.
\]

Perform the following steps for \(i = 1, 2, \ldots\).
2. Introduce the scores $t_i$ and the weights $\omega_i$,

$$t_i = E'_{i-1}\omega_i, \quad \omega_i = C[E_{i-1}, f_{i-1}].$$

3. Determine the loadings $p_i, q_i$ by least squares,

$$p_i = C[E_{i-1}, t_i]D[t_i]^{-1}, \quad q_i = C[f_{i-1}, t_i]D[t_i]^{-1}.$$

4. Find new residuals,

$$E_i = E_{i-1} - p_i t'_i \quad f_i = f_{i-1} - q_i t_i.$$

Note that in Step 3, $\omega_0 = C[X, y]$. The crucial part of the algorithm of PLS is Step 2, where the latent variable (score $t$) is formulated. The scores are defined as a linear combination of the $X$-residuals from the previous step. The choice of the weights can be justified by the property that the sample covariance between $y$ and $t_i$ is maximum in each step (Höskuldson, 1988; Stone and Brooks, 1990). This original algorithm implies that the scores are orthogonal.

Another commonly used PLS algorithm was formulated by Martens (1985) and gives orthogonal loadings. In this algorithm, $\omega_i = C[E_{i-1}, y]$ is used in Step 2, and then $E_{i-1}, f_{i-1}$ are replaced with $X, y$, respectively. As proved by Helland (1988), these two algorithms are equivalent, i.e. lead to the same predictor of $y$. However, it has turned out that the latter algorithm provides a better basis for several mathematical results of PLS. There are other versions of the PLS algorithm, for example, SIMPLS by de Jong (1993) and kernel PLS by Rännar et al. (1994), which are theoretically equivalent. For a review of the different versions of the PLS algorithm and their numerical properties, e.g. speed, we refer to Andersson (2009).

A few exact mathematical and theoretical results of the sample versions PLS are collected below.

(i) The weights $\omega_a$ satisfy the recursive relation:

$$\omega_{a+1} = s - SG_a(G'_a S G_a)^{-1} G'_a s,$$

where $s$ is the sample covariance between $X$ and $y$ and $S$ is the sample variance of $X$, and $G_a$ is any matrix spanning the column space $\zeta(\omega_1 : \omega_2 : \cdots : \omega_a)$.

(ii) The weights $\omega_1, \omega_2, \cdots, \omega_a$ span the same column space as the Krylov sequence $s, S s, \cdots, S^{a-1} s$.

(iii) The weights $\omega_i$ are orthogonal, i.e. $\omega'_i \omega_j = 0, i \neq j$.

(iv) At step $a$, PLS produces a predictor as

$$\hat{y} = \bar{y} + b'_a (X - \bar{X}), \quad b_a = G_a(G'_a S G_a)^{-1} G'_a s.$$
(v) PLS will always give a higher coefficient of determination, $R^2$, than principal component regression.

(vi) PLS shrinks in the sense that

$$|b_1| \leq |b_2| \leq \cdots \leq |b_p|.$$ 

Detailed proofs for Statement (i) - (iv) are given in Helland (1988). Statement (ii) has also been proved by Manne (1987) in a different way. von Rosen (1994) gave simple proofs of Statement (i) - (iii) applying a vector space operation. Statement (v) is pointed out by de Jong (1993) and in de Jong (1995), Statement (vi) is shown.

One main criticism of PLS from most statisticians is that there does not seem to be any well-defined probability model behind PLS. In what situation then will PLS perform well? Why is PLS useful? There are several important clarifying works, especially that by Frank (1987), who used the maximum of the covariance to describe PLS, and that by Stone and Brooks (1990), mentioned earlier, who tied OLS, PCR and PLS together under a relatively well-justified umbrella, i.e. continuum regression. Helland (1988, 1990) was the first to formulate a population version of PLS, i.e. a parametric version of PLS, and derived several theoretical results.

In practice, the algorithm of PLS presented above has to stop at some point. However, it is not easy to decide when it is to stop. Usually, cross-validation has been applied. Therefore, it can be difficult to evaluate PLS because the contribution made by performing cross-validation is unclear.

### 2.2.2 The population version of PLS

In order to cast PLS in the parametric statistical modelling framework, it is necessary to extend the algorithm of PLS in a population version, which assumes infinity of data, i.e. $n \to \infty$. Consequently, the sample mean $\bar{X}$, $\bar{y}$ and the sample covariance $S$, $s$ will be replaced with $\mu_x$, $\mu_y$, $\Sigma$, $\omega$, the parameters formulated in the model in (2.1). Then the algorithm for the population version of PLS, which was given in Helland (1990), using variables instead of data, runs as follows.

1. Define the starting values for the $x$ residuals $e_i$ and the $y$ residuals $f_i$,

$$e_0 = x - \mu_x, \quad f_0 = y - \mu_y.$$ 

Perform the following steps for $i = 1, 2, \ldots$.

2. Introduce the scores $t_i$ and the weights $\omega_i$,

$$t_i = e'_{i-1} \omega_i, \quad \omega_i = C[e_{i-1}, y].$$

3. Determine the $x$ loadings ($p_i$) and the $y$ loading ($q_i$) by least squares,

$$p_i = C[e_{i-1}, t_i] / D[t_i], \quad q_i = C[f_{i-1}, t_i] / D[t_i].$$
4. Find new residuals

\[ e_i = e_{i-1} - p_i t_i, \quad f_i = f_{i-1} - q_i t_i. \]

At each step \( a \), two linear representations are obtained,

\[ x = \mu_x + p_1 t_1 + p_2 t_2 + \cdots + p_a t_a + e_a \]
\[ y = \mu_y + q_1 t_1 + q_2 t_2 + \cdots + q_a t_a + f_a. \]

The theoretical results listed in the previous section are still valid for in the population version, and we will only mention a few which are the most crucial ones for the rest of the thesis.

(i) \( \omega_{a+1} = (I - \Sigma G_a (G'_a \Sigma G_a)^{-1} G_a) \omega. \)

(ii) \( \hat{y}_{a, PLS} = \hat{\beta}_a (x - \mu_x) + \mu_y, \) with \( \hat{\beta}_a = G_a (G'_a \Sigma G_a)^{-1} G'_a \omega. \)

(iii) \( \zeta(G_a) = \zeta(\omega_1 : \omega_2 : \cdots : \omega_a) = \zeta(\omega : \Sigma \omega : \cdots : \Sigma^{a-1} \omega). \)

In addition, there are some properties which only hold in the population version of PLS. Let \( m \) be a maximal dimension of the sequences \( \omega_1, \omega_2, \cdots, \omega_i \); i.e. \( m \) is the first integer such that \( \omega_{m+1} = 0 \). Then the following statement are true.

(i) The column space of \( G_m \), i.e. \( \zeta(G_m) \), is the smallest \( \Sigma \)-invariant space, i.e. \( \zeta(\Sigma G_m) \subseteq \zeta(G_m) \).

(ii) \( \zeta(G_m) \) is the smallest space which includes \( \Sigma^{-1} \omega \).

(iii) If PLS stops naturally, i.e. \( \omega_{m+1} = 0 \), the theoretical PLS coefficient \( \beta_{m, PLS} \) will equal \( \Sigma^{-1} \omega \).

Furthermore, Statement (i) is equivalent to

\[ \zeta(\omega_1 : \omega_2 : \cdots : \omega_i) = \zeta(\omega : \Sigma \omega : \cdots : \Sigma^{i-1} \omega) \subseteq \zeta(\omega : \Sigma \omega : \cdots : \Sigma^{m-1} \omega), \]

which is valid for all \( i \), such that \( m \leq i \leq p \). Statement (i) and (ii) together imply that PLS generates an invariant subspace with an orthogonal basis which is included in \( \zeta(\Sigma) \). Helland (1990) and von Rosen (1994) gave different proofs of the above results.

### 2.2.3 PLS, relevant components and the envelope

The notation of relevant components was introduced by Næs and Martens (1985) and then used by Helland (1990, 1992), and Næs and Helland (1993), among others. If the spectral decomposition of \( \Sigma \) is \( \Sigma = \sum_{i=1}^{p} \lambda_i u_i u'_i \), then it is assumed that there exists an ordering of terms in the decomposition and an integer \( r < p \), such that \( u'_i \omega = 0, r < i < p \). These \( u_i \) are called relevant eigenvectors. The corresponding principal component scores \( u'_i (x - \mu_x) \) are called the relevant components for the prediction of \( y \).
As mentioned earlier, the population version of PLS will generate an invariant space $G_m$ which satisfies $\zeta(\Sigma G_m) \subseteq \zeta(G_m)$ and $\zeta(\Sigma^{-1} \omega) \subseteq \zeta(G_m)$. The dimension of the invariant space $m$ is equal to $r$, the number of relevant components in $x$ for prediction. The space of $G_m$ is also spanned by the relevant eigenvectors $u_i$. In particular, $\omega$ is supposed to belong to that space.

Envelopes (Cook et al., 2010, 2013; Schott, 2013) are a recently proposed general methodology for model reduction in prediction problems. Let $y = \alpha + \beta' x + \varepsilon$, where $x$ is a random predictor with the dimension $p$, and $\varepsilon$ is the random error and is uncorrelated with $x$. An envelope arises by parameterizing the regression model in terms of the smallest subspace $R$ under the following constraints: let $P_R$ be the projection onto $R$ and $Q_R = I - P_R$, then $P_R$ should be uncorrelated with $Q_R$, and $y$ should be conditionally uncorrelated with $Q_R$ given $P_R$, according to Cook et al. (2010). Based on the algebraic characterization of the envelopes, the linear combination of $P_R x$ is of the form $u' x$, where $u$ is some eigenvectors of the variance of $x$, i.e. $\Sigma$. The minimal set of such eigenvectors of $u$ is the relevant eigenvector, which is in agreement with the nomenclature in Næs and Helland (1993).

It has been shown in Cook et al. (2013) that a proper version of the envelope model is identical to the population version of the PLS model. This leads to the finding of estimators in the regression by means of the maximum likelihood approach using the envelope model. It is argued that a likelihood-based envelope estimator is less sensitive to the number of PLS components selected and that it outperforms PLS in both prediction and estimation. The connection between PLS and the envelope has given a new insight into PLS.

### 2.2.4 Other PLS-related techniques

PLS techniques have been modified to fit various situations. Sparse PLS (Lê Cao et al., 2008; Chun and Keles, 2010) is one version of PLS which combines variable selection and prediction modelling. PLS has also been found useful in classification problems, where it is called partial least-squares discriminant analysis (PLS-DA), see Barker and Rayens (2003). PLS-DA also has a sparse version, as presented in Lee et al. (2013). Multi-block PLS (Berglund et al., 1999; Næs et al., 2011) is sequential PLS for studying the relations between several blocks of data. There are some versions of PLS which have been proposed for handling grouped data, such as least squares PLS (Jørgensen et al., 2007) and sequential and orthogonalized PLS (Næs et al., 2011). Most of these modified PLS approaches are defined algorithmically.

### 2.3 Regularization methods

There is a class of methods, often referred to as regularization methods or shrinkage methods, which are designed to deal with collinearity problems. Among others, ridge regression (RR), the least absolute shrinkage and selection operator (lasso), principal component regression (PCR) and partial least squares regression (PLS) are the most popular ones and have usually worked
well in prediction.

The idea behind RR is to stabilize the regression coefficient by adding a constant, $k$, to the diagonal of the matrix $XX'$. RR has existed for quite a while and was being implement even before the seminal work by Hoerl and Kennard (1970) popularized the technique. Note that in this subsection, $X$ is considered to be centred, i.e. $E[X] = 0$. There are many ways of determining the ridge constant $k$, each of them corresponding to a specific type of ridge regression; for a review see Brown (1994). Nowadays, the common way to understand RR is by solving a least squares criterion together with a constraint. The lasso, which was proposed by Tibshirani (1996), is defined in a similar way as RR, i.e. solving a least squares criterion with a subtle different constraint. The nature of the constraint forces some of the coefficients to be exactly zero. Consequently, the lasso combines variable selection and shrinkage, which is considered as an appealing feature. The lasso method has its own generalizations, which have been summarized in Tibshirani (2011).

PCR has figured in the statistical literature for a while, with little usage before being promoted in the chemometric field. Its popularization in chemometrics was caused by the development of PLS. PLS was first presented in an algorithmic form as a modification of the NIPALS algorithm by H. Wold (Wold, 1966), for the purpose of computing principal components. Later, as noted before, S. Wold and H. Martens (Wold et al., 1983, Martens, 1985) established it as a regression method, still only in an algorithmic way and defined without any statistical model assumption. Since then, PLS has been frequently used in the world of chemometrics.

The idea behind PCR and PLS is to find a few linear combinations of the original predictors, usually defined as components (or factors), and then directly regress the response on these components. In PCR and PLS the components are formed differently. The components in PCR are formed via the eigenvectors of the covariance matrix of the predictors; i.e. the components in PCR only depend on the predictors. This has been regarded with some scepticism, since even though the components are chosen in such a way that they will explain the predictors in the best possible way, there is no guarantee that these components will be pertinent to predicting the response as well. The components in PLS are obtained by maximizing the covariance between the response and all the possible linear functions of the predictors.

In the following subsections, we will review the popular regularization methods with respect to numerical formulations, shrinkage properties, and linkages and comparisons among the regularization methods.

### 2.3.1 Numerical approaches

The term regularized emanates from the method of regularization used in approximation theory (Brown, 1993). Therefore, it is worth considering all the methods from a numerical point of view. In my opinion, the justification for using the methods is quite clear if the aim is to solve a linear system.

The basic solution for a linear system is found by minimizing the quadratic
with respect to $X$ over a proper subset, $\mathbb{R}^p$. If $X$ is collinearly structured and ill-conditioned, the straightforward solution for (2.3) becomes sensitive to data values. Therefore, one may put constraints on the solution, which is one kind of regularization. Roughly speaking, regularization is a technique for transforming a poorly conditioned problem into a stable one (Golub and Van Loan, 1996).

Ridge regression is the solution obtained by minimizing
\[
\min_{\beta} \| y - X' \beta \|^2_2 + \lambda \| \beta \|^2_2.
\]
Since $X$ is ill-conditioned, the solution $\| \hat{\beta} \|^2_2$ may become quite large. The large scale of the solution could be considered as the reason for the bad performance. Therefore, ridge regression includes the Euclidean norm (also called the $L_2$-distance); i.e. $\lambda \| \beta \|^2_2$ acts as a penalty term which leads to restrictions on the scale of the solution.

Instead of using the Euclidean norm as the penalty, the lasso uses the Manhattan norm (the $L_1$-distance) as the penalty; i.e. the lasso is the solution obtained by minimizing:
\[
\min_{\beta} \| y - X' \beta \|^2_2 + \lambda \| \beta \|^1_1.
\]

Another possible way to constrain the parameters would be to solve
\[
\min_{V' \beta = \gamma} \| y - X' \beta \|^2_2 \approx \min_{\gamma} \| y - X' V \gamma \|,
\] (2.4)
where $V$ is a matrix with orthogonal columns. The matrix $V' \beta$ can be considered as transforming the solution $\beta$ into a lower dimensional space.

PCR can be obtained by (2.4) using truncated singular value decomposition (truncated SVD). SVD states that any matrix $A_{p \times q}$ can be factorized as
\[
A = U D V',
\]
where $D = (D_r, 0)$, $D_r = diag(\sqrt{\lambda_1}, \sqrt{\lambda_2}, \ldots, \sqrt{\lambda_r})$, $\sqrt{\lambda_i}$ are the singular values, $r = rank(A)$, and $U$ and $V$ are orthogonal matrices. Truncated SVD uses the largest $k$ singular values in $D_k$ to approximate $A$, such as
\[
A \approx U_k D_k V_k',
\]
with $U = (U_k, U_{\perp})$, where $U_{\perp}$ is a $p \times (p-k)$ matrix such that $U$ is orthogonal and $V = (V_k, V_{\perp})$. Therefore, if there is a linear system,
\[
\min_{\beta} \| X y - X X' \beta \|^2_2,
\]
which needs to be solved, we use truncated SVD first and thus $XX' = U_kD_kU_k'$. The $U_k$ is used as a transformation matrix such that $U_k'\beta = \gamma$. Therefore, the linear system can be reformulated as

$$
\min_\gamma \|Xy - U_kD_kU_k'U_k\gamma\|^2_2
= \min_\gamma \|U_k'Xy - D_kI_k\gamma\|^2_2 + U'Xy.
$$

The solution to the linear system equals

$$
\hat{\gamma} = \begin{pmatrix} u_1'Xy/\lambda_1 \\ u_2'Xy/\lambda_2 \\ \vdots \\ u_k'Xy/\lambda_k \end{pmatrix}, \quad \hat{\beta} = U\hat{\gamma} = \sum_{i=1}^k \frac{u_i'Xy}{\lambda_k} u_i,
$$

where $\hat{\beta}$ mathematically equals the PCR solution.

PLS and Lanczos bidiagonalization (LBD) are equivalent mathematically (Eldén, 2003). The LBD procedure generates a series of matrices, $R_k = (r_1, \cdots r_k)$, $Q_k = (q_1, \cdots, q_k)$ and

$$
Z_k = \begin{pmatrix} \alpha_1 & \gamma_1 \\ \alpha_2 & \gamma_2 \\ \vdots & \vdots \\ \alpha_{k-1} & \gamma_{k-1} \\ \gamma_k \end{pmatrix},
$$

which satisfy $X'R_k = Q_kZ_k$. Subsequently, $R_k$ and $Q_k$ consist of orthogonal columns and span the Krylov structured spaces.

$$
\zeta(R_k) = \zeta(Xy, (XX')(Xy), \cdots, (XX')^{k-1}(Xy)),
$$

$$
\zeta(Q_k) = \zeta(X'R_k, (X'X)(X'Xy), \cdots, (X'X)^{k-1}(X'Xy)).
$$

Accordingly, if we want to compute the solution for (2.3), LBD provides a natural transformation matrix, $R_k$, such that $R_k'\beta = \gamma$. Then the solution $\gamma$ can be obtained by solving

$$
\min_\gamma \|y - X'R_k\gamma\|^2_2 = \min_\gamma \|y - Q_kZ_k\gamma\|^2_2
= \min_\gamma \|Q_k'y - Z_k\gamma\|^2_2 + \|Q_k'y\|^2.
$$

Thus the solution is given by

$$
\hat{\gamma} = Z_k^{-1}Q_k'y, \quad \hat{\beta} = R_kZ_k^{-1}Q_k'y.
$$

It can be shown that the above $\hat{\beta}$ is mathematically equivalent to the derivation of estimators via the sample version of PLS.
2.3.2 Shrinkage property

Based on the estimators derived from the numerical approaches, it is convenient to explore the shrinkage property of the regularized estimators. Frank and Friedman (1993) defined the “shrinkage factor” concept to compare the shrinkage behaviour of the methods. The general proposed form of the estimators is

$$\hat{\beta} = \sum_{j=1}^{r} f(\lambda_j) \hat{\alpha}_j u_j,$$

with

$$\hat{\alpha}_j = \frac{1}{\lambda_i} u_j' X y, \quad \sum_{j=1}^{r} \left( \frac{1}{\lambda_i} u_j u_j' \right) = X X' ,$$

where $r$ is the rank of $X$ and $f(\lambda_j)$ are called shrinkage factors. For the MLE, $f(\lambda_j) = 1$. If $f(\lambda_j) < 1$, this will lead to a reduction in the variance of $\hat{\beta}$, although it may introduce a bias as well. It is hoped that any increase in the bias will be small compared to the decrease in the variance, so that the shrinkage will be beneficial. In ridge regression, the shrinkage factor $f(\lambda_j) = \lambda_j / (\lambda_j + \lambda)$, which is always smaller than 1. For principal component regression, $f(\lambda_j) = 1$ if the $j$th component is included. Otherwise, $f(\lambda_j) = 0$.

The shrinkage property for PLS is peculiar (Butler and Denham, 2000). The shrinkage factor $f(\lambda_j)$ is not always smaller than 1. The component corresponding to the smallest eigenvalue can always be shrunk. The shrinkage factor $f(\lambda_1) > 1$ if the number of components in PLS is odd, and the shrinkage factor $f(\lambda_j) < 1$ if the number of components in PLS is even. Björkström (2010) showed that the peculiar pattern of alternating shrinkage and inflation is not unique for PLS. For a review of the shrinkage properties of PLS, we refer to Krämer (2007).

In summary, RR shrinks all directions, but has a greater effect on the low-variance direction. PCR only shrinks the first $a$ high variance and discards the rest. PLS shrinks the low-variance directions, but, peculiarly enough, inflates some of the high-variance directions as well.

2.3.3 Linkage among the regularization methods

One main stream of the discussion on regularization methods concerns the linkage among them. Among others, Stone and Brooks (1990) introduced continuum regression (CR), where OLS, PCR and PLS all naturally appear as special cases. CR is formulated as choosing latent components by maximizing

$$T(\gamma, c) = (c' X' y) (c' X' X c)^{\gamma-1},$$

where $c$ is a vector with $|c| = 1$. After finding this $c$, the regression coefficients $\beta$ in the prediction are constructed by performing simple linear regression, $y$ on $X c$. Here, $\gamma = 0$ gives OLS in one step by maximizing the correlation, $\gamma = 2$ gives PLS via maximizing the covariance, and $\gamma = \infty$ corresponds to
PCR by maximizing the variance. Furthermore, the relationship between RR and CR was pointed out by Sundberg (1993), who explained CR with one component differs from RR only by a scalar factor, which is a function of the ridge constant \( \lambda \). Any CR regressor with \( \gamma \) between 0 and 1 is in fact a ridge regressor. As implemented by de Jong and Farebrother (1994), this correspondence can be extended to \( \gamma > 1 \) in CR by using a negative ridge constant.

From a conceptual point of view, CR is very attractive in that it ties OLS, PCR, PLS and RR together in a unified framework and enhances the understanding of various methods and their intimate relationship. However, the methodology suffers from both a heavy computation burden (for example, due to its use of cross-validation to estimate \( \gamma \) and the number of latent components) and some inferential opaqueness (Brown, 1993). It should be mentioned that Brooks and Stone (1994) also proposed a multivariate version of CR, joint continuum regression. Both the univariate and the multivariate versions have so far not been used much in applications.

2.3.4 Comparisons

Which method (when comparing among RR, the lasso, PLS, and PCR) is the best one? There is consensus that typically all these methods are approximately equivalent and possess a relatively better prediction ability than OLS and variable subset selection (VSS) for collinear data.

Due to the fact that PLS is only algorithmically defined, it is not easy to draw any firm conclusions by comparing all the methods under a general model set-up. However, there are several empirical studies documented in the literature. Hoerl et al. (1975), Gibbons (1981) and Muniz (2009) compared different types of RR, Lawless (1976) focused on RR and PCR, Dempster et al. (1977) compared many shrinkage-type estimators, and Garthwaite (1994) conducted a comparison between PCR, PLS and VSS, etc. This list could have been made much longer, but we will below only provide details of a few research studies.

Among others, Frank and Friedman (1993) conducted an extensive simulation study and concluded that RR is generally preferable to VSS, PLS and PCR, but that the superiority of RR over PLS and PCR was so slight that “one would not sacrifice much average accuracy over a lifetime by using one of them to the exclusion of the other two”. Tibshirani (1993) compared the performance of the lasso with that of RR and VSS, and concluded that, when there are a ‘large number of small effects’, RR is best, followed by the lasso and then VSS; when there are a ‘small to moderate number of moderate-sized effects’, the lasso performs best, followed by RR and VSS; and VSS performs much better than both RR and the lasso with a ‘small number of large effects’.

It is worth mentioning that Helland and Almøy (1994), who presented an asymptotic result, compared PLS and PCR using a multinormal model when only a few components were relevant, i.e. when only some of the population version of principal components in the explanatory space were correlated with
the dependent variable. Their conclusion is that the difference between PCR and PLS in most cases is relatively small. PCR performs better when the irrelevant eigenvalues are relatively small or relatively large. For intermediate irrelevant eigenvalues, PLS performs better. In practice, PLS may be preferable, since large irrelevant eigenvalues rarely exist and the difference is very small for small irrelevant eigenvalues. Almøy (1996) carried out a simulation study which confirmed the above conclusions.

In summary, RR, the lasso, PLS and PCR perform similarly. RR is preferable in some cases only to a very limited extent. The lasso can be used when both shrinkage and variable selection are needed.

2.3.5 Other shrinkage methods

In the literature, many other shrinkage methods are suggested, which are used relatively less frequently than the above-mentioned ones: intermediate least squares regression (Frank, 1987), James-Stein shrinkage (James and Stein, 1961), latent root regression (Webster, 1974), reduced rank regression (Andersson, 1958; Izenman, 1975; Reinsel and Velu, 1998), least angle regression (Efron et al., 2004), various Bayes-Stein-type estimators (Zellner, 1972) and other Bayes methods. For a review, see Dempster (1977).

2.4 The growth curve model and its extensions

It appears that, when developing our two-step approach, the classical growth curve model and its extensions have played a key role. Therefore, in the subsequent subsections, these models are presented in some detail.

2.4.1 The growth curve model

The growth curve model was proposed by Potthoff and Roy (1964) and has many important applications within medicine, social sciences, etc.

Let \( \mathbf{X} : p \times n \), \( \mathbf{A} : p \times q, q \leq p \), \( \mathbf{B} : q \times k \), \( \mathbf{C} : k \times n \), \( r(\mathbf{C}) + p \leq n \), where \( \mathbf{\Sigma} : p \times p \) is positive definite. Then,

\[
\mathbf{X} = \mathbf{ABC} + \mathbf{E} \tag{2.6}
\]

defines the growth curve model, where the columns of \( \mathbf{E} \) are assumed to be independently distributed as a multivariate normal distribution with mean 0 and a positive definite dispersion matrix, \( \mathbf{\Sigma} \), i.e. \( \mathbf{E} \sim N_{p,n}(0, \mathbf{\Sigma}, \mathbf{I}_n) \). The matrix \( \mathbf{C} \) is often called a between-individuals design matrix and is precisely the same design matrix as that used in the univariate linear model. The matrix \( \mathbf{A} \) is often called a within-individuals design matrix. Both the matrices \( \mathbf{A} \) and \( \mathbf{C} \) are known, whereas the matrices \( \mathbf{B} \) and \( \mathbf{\Sigma} \) are unknown parameter matrices.

The dental data set of Potthoff and Roy (1964) will be used to illustrate the growth curve model. The data were obtained through dental measurements performed on 11 girls and 16 boys at four different ages \( t_1 = 8, t_2 = 10, \ldots \)
Figure 1: The distance from the centre of the pituitary to the pteryomaxillary fissure in girls (○) and boys (+) at the ages of 8, 10, 12 and 14. The growth profiles for girls (−) and boys (−−) are assumed to be linear in time.

t_3 = 12, t_4 = 14). Each measurement is the distance from the centre of the pituitary to the pteryomaxillary fissure. These data are plotted in Figure 1.

Let us consider a case where the mean of the distribution for each treatment group is supposed to be linear. In that case the \( \mu_i \) for group \( j \) at time \( t_i \) is given by

\[
\mu_j = \beta_{1j} + \beta_{2j}t_i \quad j = 1, 2, i = 1, 2, 3, 4.
\]

The size of the observation matrix \( X \) is 4 \times 27, and the first 11 columns correspond to measurements on girls, while the last 16 columns correspond to measurements on boys. The between-individuals design matrix \( C : 2 \times 27 \) equals

\[
C = \left( 1_{11}^\prime \otimes \begin{pmatrix} 1 \\ 0 \end{pmatrix} : 1_{16}^\prime \otimes \begin{pmatrix} 0 \\ 1 \end{pmatrix} \right),
\]

where \( \otimes \) means the Kronecker product. Due to the linear mean structure, the within-individuals design matrix \( A : 4 \times 2 \) equals

\[
A^\prime = \begin{pmatrix} 1 & 1 & 1 & 1 \\ 8 & 10 & 12 & 14 \end{pmatrix}.
\]

Then the expectation of the data matrix \( X \) can be presented as \( E[X] = ABC \) and the variance \( D[X] = I \otimes \Sigma \), where \( B \) and \( \Sigma \) are unknown matrices. Thus, we may use the growth curve model \( X \sim N_{4,27}(ABC, \Sigma, I) \).

The maximum likelihood method is one of the approaches used to find estimators of the parameters in the growth curve model. The explicit maximum
likelihood estimators (MLEs) in the growth curve model have been derived by different approaches by different authors, see Kollo and von Rosen (2005).

2.4.2 The extended growth curve model

In the growth curve model, each group follows a linear growth profile. In some cases, this may not hold, as shown in Figure 2.

![Figure 2: The distance from the centre of the pituitary to the pteryomaxillary fissure in girls (○) and boys (+) at the age of 8, 10, 12 and 14. The growth profiles for girls (−) and boys (−−) are assumed to be different, as defined in the model in (2.7).](image)

For example, it may be more reasonable to assume that the means of the boys’ group follows a second degree polynomial, i.e.

\[ \mu_2 = \beta_{21} + \beta_{22}t_i + \beta_{23}t_i^2, \quad i = 1, 2, 3, 4. \]

Thus, a natural way to extend the classical growth curve model is given by:

\[
X = A_1B_1C_1 + A_2B_2C_2 + E, \quad \text{(2.7)}
\]

where

\[
A_1' = \begin{pmatrix}
1 & 1 & 1 & 1 \\
8 & 10 & 12 & 14
\end{pmatrix}, \quad C_1 = \begin{pmatrix}
1'_{11} \otimes \begin{pmatrix}
1 \\
0
\end{pmatrix} & 1'_{16} \otimes \begin{pmatrix}
0 \\
1
\end{pmatrix}
\end{pmatrix},
\]

\[
A_2' = \begin{pmatrix}
8^2 & 10^2 & 12^2 & 14^2
\end{pmatrix}, \quad C_2 = \begin{pmatrix}
0'_{11} & 1'_{16}
\end{pmatrix}.
\]

The matrices \(B_1\) and \(B_2\) are parameter matrices and \(E\) is the same normally distributed error matrix as before, i.e. \(E \sim N(0, \Sigma, I)\). The model in (2.7)
assumes that for both girls and boys we have a linear structure, but additionally that for the boys there exists a second-order degree polynomial structure. The model in (2.7) satisfies the condition $\zeta(C'_2) \subseteq \zeta(C'_1)$, which is a crucial relation in order to obtain explicit estimators.

The model in (2.7) can be formulated in a more general form. Let $X: p \times n$, $A_i: p \times q_i$, $B_i: q_i \times k_i$, $C_i: k_i \times n$, $r(C_i) + p \leq n$, $i = 1, 2, \ldots, m$, $\zeta(C'_i) \subseteq \zeta(C'_{i-1})$, $i = 2, 3, \ldots, m$ and $\Sigma: p \times p$ be positive definite. Then,

$$X = \sum_{i=1}^{m} A_iB_iC_i + E,$$

(2.8)

where $E \sim N_{p,n}(0, \Sigma, I)$. The model in (2.8) is called an extended growth curve model. The model in (2.7) is a special case of that in (2.8) when $m = 2$. An exhaustive description of the extended growth curve model can be found in Kollo and von Rosen (2005).

A crucial condition for obtaining explicit maximum likelihood estimations (MLEs) of the extended growth curve model is $\zeta(C'_i) \subseteq \zeta(C'_{i-1})$, $i = 2, 3, \ldots, m$, $C_0 = I$. It may be worth mentioning that the subspace conditions $\zeta(C'_i) \subseteq \zeta(C'_{i-1})$ may be replaced with $\zeta(A_i) \subseteq \zeta(A_{i-1})$. How to obtain the MLEs in the extended growth curve model is described in Kollo and von Rosen (2005).
3 Summary of papers

3.1 A real data set which has inspired Papers I-IV

3.1.1 Data description

Silage sample data were prepared for experimental purposes at the Swedish University of Agricultural Sciences during 2002-2006, and these data covered a total of 762 silage samples from 15 different experiments. The samples were ensiled in mini-silos of varying size for a minimum of 60 days. After ensiling, silage juice was obtained with a hydraulic press, and was later prepared for chemical analysis, among other types of analysis.

Reference analyses were performed of ten soluble compounds in the silage, i.e. ammonia nitrogen, lactic acid, acetic acid, propionic acid, butyric acid, total volatile fatty acids, succinic acid, butanediol, ethanol and water-soluble carbohydrates (WSC or sucrose). High-performance liquid chromatography (HPLC) was used for the analysis of lactic acid, acetic acid, propionic acid, butyric acid, total volatile fatty acids, succinic acid, 2,3-butanediol and ethanol. The HPLC system consisted of a Hewlett Packard Series 1050 pump, a Marathon (Spark Holland BV, the Netherlands) auto-sampler with a 20 µl loop and an ERC-7510 (ERMA Inc., Japan) RI-detector. A 300×7.8 mm stainless-steel column, packed with ReproGel H, and a pre-column packed with the same material were used. The mobile phase consisted of 0.005 M sulphuric acid and the flow rate was 0.8 ml/min. The water-soluble carbohydrates were analyzed using an enzymatic method (Udén, 2006) applied to dried samples, and the concentration in the silage juice was estimated from the water content of the silage. The ammonia nitrogen concentrations were analyzed using the phenol-hypochlorite and ninhydrin colorimetric assays adapted to continuous-flow analysis (Broderick and Kang, 1980; Broderick, 1987). The unit used for all the reference analyses was grams per litre.

All the samples were analyzed two to four years after the time of the reference analysis using Fourier transform infrared (FTIR) analysis. The FTIR instrument used had originally been designed for routine milk analysis (Milcoscan FT120, Foss Electric A/S, Hillerød, Denmark), but was modified to allow the liquids to be pumped directly into the measurement cell. The instrument measured mid-infrared spectra from 999 cm$^{-1}$ to 4996 cm$^{-1}$. The spectra were recorded at 1,037 wave numbers at intervals of 3.858 cm$^{-1}$. Thereafter, the spectra with the observation means subtracted were used in calibration.

The aim of the analysis was to be able to utilize data on a small number of samples where both reference values and FTIR data are available to develop a predictive model for the chemical content in a silage sample based only on the FTIR spectra.
3.1.2 A short background to the interpretation of the spectra

According to Beer’s law (see Cross, 1969), spectral peak absorbance values are linearly related to concentrations of soluble compounds in a solution. As illustrated in Figure 3, the higher the concentration of sucrose is, the bigger are the absorbance values in the spectra. There are two peaks in the spectra (Peak 1 at a wave number around 1158 cm$^{-1}$ and Peak 2 at a wave number around 1019 cm$^{-1}$) which can be considered as the two main underlying components which are directly connected to the effect of sucrose. This means that, based on the spectra of the pure solution, a model for the concentration of sucrose can be built using only these two components (peaks).

The overall spectra of the silage reveal, however, mixtures of ten known soluble compounds and a number of unknown ones. An examination of the spectra of pure solutions of a few common compounds (Figure 4) shows considerable overlap. Nevertheless, the complex spectrum of multiple compounds can still be said to correspond to the sum of all the individual spectra. This means that in mixtures, no compound will correspond to any single peak, as each peak is influenced by all the compounds. Therefore, when building a model for an individual compound, it is necessary to extract a number of important signals from the mixture. For example, if we intend to build a model for sucrose, the original Peak 2 at a wave number around 1019 cm$^{-1}$ may still be useful, but it is also influenced by lactate and ethanol. Accordingly, we need to consider other regions where, for example, lactate is dominant, in order to correct Peak 2 for the effect of lactate. In reality, multiple peaks are used, some of which are linked to peaks of the compound of interest and others to spectral regions of interfering compounds.

Figure 3: The spectra of two solutions of sucrose. The upper curve is the spectrum of sucrose with a concentration of 200g/L, while the lower curve is the spectrum of sucrose with a concentration of 100g/L.
Figure 4: The spectra of the mixture and the pure solutions. The black solid curve is the spectrum of the mixture solution, while the coloured dashed curves are the spectra of pure compounds, i.e. sucrose, lactate, acetate and ethanol.

3.1.3 How the data inspired the papers

The multiple characteristics of the data, for example their collinearity and group effects, inspired this thesis. First of all, the data consist of collinear variables. The aim is to build a model for soluble compounds using the spectra. Each wave number in the spectrum is an explanatory variable. Naturally, adjacent wave numbers are correlated. Paper I is devoted to an investigation of the performance of the most popular regression methods suitable for handling collinear data.

The dominant method for such data in the chemometric field is PLS, which is mainly algorithmically based and lacks a proper statistical model. Therefore, the intention of Paper II is to give an explanation of PLS in a statistical context where a new two-step method is introduced.

There are group effects in the silage data, since the data originates from different experiments. In some experiments, the data shared the same structure, i.e. concerned the same kind of soluble compounds, but had different mean levels. For example, it is natural that the mean level of ethanol should be high in one experiment, but low in another due to the silage-type in question. An illustration of the spectra from two experiments is provided in Figure 5, where each colour represents an experiment (group). The two groups of spectra show peaks in the same region of wave numbers, but at two distinct levels. Paper III is devoted to such grouped data, i.e. data with different mean levels and the two-step method is extended in this paper to comprise group effects.

The samples in some experiments had been exposed to special treatments,
Figure 5: *Spectra of the silage samples from two different experiments.* Each experiment is represented by one colour. The spectra have the same pattern, but two different levels, which indicates that the means of the two groups follow the same structure, but have different levels.

Figure 6: *Spectra of the silage samples from two different experiments.* Each experiment is represented by one colour. There is an extra peak in one experiment, as indicated by the dashed line, which implies that the mean structure of one group is more complicated than that of the other one.
as a result of which the samples in these experiments comprised a special soluble component (e.g. ammonia) which was absent or had very low values in other experiments. The spectra for two such experiments are shown in Figure 6 where the two groups of spectra share several peaks in some regions of wave numbers. However, in one experiment there is an extra peak which does not exist in the other one. Thus, the structure of the data in some experiments is more complicated than that in the rest. In fact, the mean structure from one group is nested in another, which is called a nested group effect. Therefore, Paper IV is devoted to prediction when this type of structure exists.

3.2 The linkage between the papers of the thesis

Figure 7 represents a flowchart showing how the silage data inspired the papers, as well as the connections between the papers. In comparison with the other papers, Paper I is relatively independent of the others and served as an overview of the commonly used methods for collinear data. A summary of the results of Paper I is presented in Section 3.3. The contents of Paper II-IV are closely connected. As shown in Figure 7, Paper II-IV are linked in that the data structures dealt with therein become increasingly complex with each successive paper. Some of the contents of Paper II-IV overlap. For example, the idea of the two-step method is proposed in Paper II, but is further exploited in Paper III through the addition of a new explanation. The intention of the following subsections is to summarise the contents in a logical way, which does not strictly follow the order of the papers. Sections 3.4-3.8 treat the development of the two-step method. The main idea of the two-step method is presented in Section 3.4 based on Paper II and III. The discussion of the connections between the two-step method and PLS in Section 3.5 is based on Paper II and III. Section 3.6, 3.7 and 3.8 deal with results presented in Paper II, III and IV.

Figure 7: A schematic flowchart for the relation between the silage data and the papers of the thesis and the linkage between the papers.
3.3 An empirical study of popular shrinkage methods

In Paper I, we investigated the robustness of the popular shrinkage methods, i.e. ridge regression, the lasso, partial least squares regression and principal component regression, together with OLS and variable subset selection (VSS), through their applications on a real data set.

A part of the silage data set was selected which consisted of data on a total of 630 grass-clover silage samples from 13 different experiments. Reference analyses of ten soluble compounds in the silage had previously been made for these samples. The spectra from 999 to 1585 cm$^{-1}$ for the same samples were used.

The structure of the silage data is rather complex. Besides collinearity, there are several features which are rarely discussed. Firstly, the data originate from several experiments. The differences between the experiments, for example concerning the silage type and the experimental conditions, are unknown or too complicated to be summarized. Furthermore, multi-responses are, naturally, present in the data. The covariance structure of the multi-response variables varied to a large extent among the experiments. The influence of the covariance between the responses on the prediction is unknown.

There is a lack of appropriate shrinkage methods for such data, i.e. data characterised by group (experiment) effects and multi-responses. Therefore, deliberately neglecting those structural characteristics of the data, we examined the performance of the most popular shrinkage methods using uni-response models without any group information. To a certain extent, the models were misspecified. Our purpose was to investigate the robustness of the popular shrinkage methods under these misspecified models.

To evaluate the methods, two procedures were used in Paper I. In Procedure 1, the samples were randomly divided into two subsets. One subset contained 567 samples (around 90 percent of the total number of samples) and was used to build the models, and throughout the paper it is called the training set. The other 63 samples made up a subset which was used for testing the model performance and is called the test set. A 10-fold cross-validation was employed within the training set to determine the tuning parameter for each method. The whole routine described above was repeated 100 times. In Procedure 2, in each repetition, one of the 13 experiments was saved to serve as a test set. The other 12 experiments (the training set) were used to build the model. This process was repeated 13 times with the data from each experiment used only once for validation. The tuning parameters were determined via 12-fold cross-validation within the training set, with each subset corresponding to one of the 12 experiments. In each procedure, for one response (analyte), each method was ranked by its root mean squared error for prediction. Then we assigned each calibration method a score according to its ranking. The final score for each method within each procedure was the sum of all of its scores across all the analytes, which was used as a criterion for the overall performance.

For the overall performance, as summarized in Paper I, the ranking for
Procedure 1 was:

Lasso > RR > VSS > PLS > OLS > PCR.

The ranking for Procedure 2 was:

Lasso > RR > VSS > PLS > PCR > OLS.

In conclusion, we propose the use of the lasso and RR for complex data due to the good robustness properties of these methods. Classical VSS may also be a good choice because the outcome is relatively easy to interpret. PCR showed an unexpectedly poor performance. Applying PCR with a uni-response model to multi-response data should therefore be carried out with caution.

3.4 PLS viewed using a two-step estimation approach

The population PLS predictor at step $a$ equals, if $\mu_x$ and $\mu_y$ are known,

$$\hat{y}_{a,PLS} = \omega'G'_a(G'a\Sigma G_a)^{-1}G'_a(x - \mu_x) + \mu_y.$$  \hspace{1cm} (3.1)

We have formulated the above classical PLS predictor as a two-step estimation approach, which will give us new insight into PLS. This idea, initially discussed in Paper II, is supplemented in Paper III. In the first step, with the assumption that $x$ is proportional to the covariance $\omega$, it is supposed that the following model holds, with $\varepsilon \sim N_p(0, \Sigma)$,

$$x - \mu_x = \omega \gamma + \varepsilon = \Sigma \Sigma^{-1} \omega \gamma + \varepsilon.$$  \hspace{1cm} (3.2)

The product $\Sigma \Sigma^{-1}$ is used to enable the canceling-out of $\Sigma^{-1}$ in the conditional predictor, which causes the poor performance when estimating $\Sigma$ with near-collinear data. Based on the Cayley-Hamilton theorem, $\Sigma^{-1}$ can be presented in a polynomial form, i.e. $\Sigma^{-1} = \sum_{i=1}^{p} c_i \Sigma^{-1} \approx \sum_{i=1}^{a} c_i \Sigma^{-1}$, for some $c_i$, which is a function of $\Sigma$ and $a \leq p$. One important simplification here is that, instead of considering $c_i$ as a function of $\Sigma$, we treat it as an unknown constant which, together with $\gamma$, should be estimated later. If $\Sigma$ is unknown, then $\{c_i\}$ is also unknown. If $a = p$, there is no approximation. How to determine $a$ is an open question. Consequently,

$$x - \mu_x = \Sigma \sum_{i=1}^{p} c_i \Sigma^{-1} \omega \gamma + \varepsilon \approx \sum_{i=1}^{a} \Sigma \omega \beta_i + \varepsilon = \Sigma G_a \beta + \varepsilon,$$  \hspace{1cm} (3.3)

where $\beta = (\beta_i)$ and $\beta_i = c_i \gamma$ is an unknown parameter vector. The matrix $G_a$ was defined in the presentation of the population version of PLS, i.e. $\zeta(G_a) =$
\(\zeta(\omega : \Sigma \omega : \cdots : \Sigma^{a-1} \omega)\). It may be helpful to view \(\mu_x\) as a baseline parameter rather than a population mean as defined before. In PLS applications, data which are pre-centered with the sample mean are usually used. The model in (3.3) models the mean with adjustment for the baseline. As mentioned earlier, the population version of PLS will generate an invariant space which has the property \(\zeta(\omega) \subseteq \zeta(\Sigma G_a)\). Then \(\omega = \Sigma G_a \rho\) will transform the model in (3.2) into that in (3.3). The model in (3.3) satisfies the assumption that it is a weakly singular Gauss-Markov model (see Nordström, 1985). Then a least squares predictor is given by

\[
\hat{x} = \Sigma G_a (G_a' \Sigma \Sigma^{-1} \Sigma G_a) - G_a' \Sigma \Sigma^{-1} (x - \mu_x) = \Sigma G_a (G_a' \Sigma G_a) - G_a' (x - \mu_x).
\]

In the second step,

\[
\hat{y} = \omega' \Sigma^{-1} (\hat{x} - \mu_x) + \mu_y = \omega' G_a (G_a' \Sigma G_a) - G_a' (x - \mu_x) + \mu_y
\]

is used, is identical to \(\hat{y}_{a, PLS}\) in (3.1), and is completely free of \(\Sigma^{-1}\).

So far we have established the linkage between PLS and a two-step estimation approach. Using a design matrix, \(A = \Sigma G_a\), in the first step will lead to a predictor identical to the classical PLS predictor. The choice of the design matrix was motivated by the Cayley-Hamilton theorem as an approximation of \(\Sigma^{-1}\). This gives a new insight into PLS; i.e. PLS can be viewed as a method of generating a Krylov structured space to approximate \(\Sigma^{-1}\). Moreover, it has been shown in von Rosen (1994) that when selecting a design matrix which gives the PLS predictor and also satisfies the weakly singular Gauss-Markov model, one is limited to those matrices generating \(\zeta(\Sigma G_a)\).

Another way to view the model in (3.3) is to assume \(\beta\) to be random with \(E[\beta] = 0, D[\beta] = I, C[\beta, \epsilon] = 0\) and to assume \(\mu_x\) to be the population mean. Then the model in (3.3) indicates that the dispersion of \(x\), i.e. \(\Sigma\), can be decomposed into two parts. One part is structured and belongs to the space of \(\zeta(G_a)\), which is the same space as that generated by the relevant components mentioned earlier. Consequently, the model in (3.3) follows the set-up of a mixed linear model. Thus, the problem is to estimate \(\mu_x\) and \(\Sigma\) and to predict \(\beta\). Inserting the estimators, the conditional predictor becomes

\[
\hat{y} = \omega' \Sigma^{-1} (x - \hat{\mu}_x - \hat{\Sigma} G_a \hat{\beta}) + \mu_y.
\]

3.5 A new two-step regression method

Inspired by PLS, a new two-step method is proposed for collinear and high-dimensional data. The two-step method is formulated as follows:

(i) \(x = \omega \gamma + \varepsilon, \varepsilon \sim N(0, \Sigma)\);

(ii) \(y = \omega' \Sigma^{-1} (x - \hat{\mu}_x) + \mu_y\).
The first step is the step for summarizing the information in \( x \). For collinear data, often with a relatively large number of \( x \) variables, it is fairly reasonable to assume a smaller number of ‘latent components’ that influence the explanatory variables and thus the response variable as well. As illustrated in the silage data shown in Figure 4, each peak represents some underlying components. Random effects are usually present in the explanatory variables as well, for example some measurement errors. Thus, the information is summarized in a model where the latent components are treated as fixed effects and the error is considered to be a random effect. The second step is the prediction step. A natural choice is to use a conditional predictor, since the conditional predictor is the best choice in that it has the smallest variance among unbiased predictors.

The two-step modelling idea, i.e. predicting \( y \) via some form of summarized \( x \), is not new. For example, Stone and Brooks (1990) in continuum regression used a ‘potential additional regressor’, which is based on \( x \) under some maximum criteria. Helland (1992) proposed a maximum likelihood approach using the relevant components which is based on the eigenvectors of \( \Sigma \).

A critical problem is how to estimate \( \Sigma^{-1} \) in Step 2. The collinear data will lead to the estimator of \( \Sigma \) being close to singular and then the inverse of the estimator will have a large dispersion, which results in a poor predictor. In the literature, the approaches to approximating \( \Sigma^{-1} \) are removal of the eigenvectors with small eigenvalues, i.e. shrinkage, and the use of the regularization approach \((\Sigma + \lambda I)^{-1}\), which is similar to ridge-type regression. We use a new approach inspired by the Cayley-Hamilton theorem as applied in PLS, i.e. \( \Sigma^{-1} = \sum_{i=1}^{p} c_i \Sigma^{-1} \), where \( c_i \) are called generalized traces and functions of \( \Sigma \). Thus, we have a model of the form

\[
\mathbf{x} = \Sigma \mathbf{G}_a \beta + \epsilon, \tag{3.4}
\]

in the first step. It is worth mentioning that PLS formulated in the two-step approach starts with \( x - \mu_x \), which is proportional to \( \omega \), whereas the new approach starts with \( x \) being proportional to \( \omega \). If one considers the model \( x - \mu_x = \omega \gamma + \epsilon \) to be modelling an adjusted mean by treating \( \mu_x \) as a baseline parameter, this model is essentially the same as \( x = \omega \gamma + \epsilon \), since both model the mean, with the former doing so for pre-treated data and the latter doing so for raw data. Therefore, instead of using the sample mean to approximate the baseline and estimate the adjusted mean from the model \( x - \mu_x = \omega \gamma + \epsilon \), it is fairly reasonable to simplify the model as \( x = \omega \gamma + \epsilon \) and estimate the population mean directly. If one considers the model \( x - \mu_x = \omega \gamma + \epsilon \) to be modelling the dispersion of \( x \), as discussed earlier, this model is different from the model \( x = \omega \gamma + \epsilon \), which models the mean.
3.6 The two-step method for linear prediction

3.6.1 Model

The connection between PLS and the two-step method provides a natural choice for the design matrix $A$, i.e. $\Sigma G_a$. Now, with $n$ pairs $(y_i, x_i)$, $i = 1, 2, \ldots, n$, of independent observations, the model is formulated as:

$$X = A\beta_1' + E,$$  \hspace{1cm} (3.5)

with $X: p \times n$, $A: p \times q$, $\beta: q \times 1$, and where $1_n': 1 \times n$ is a vector of $n$ 1s, $E \sim N_{p,n}(0, \Sigma, I_n)$, $A = \Sigma G_a$, $G_a$ is the Krylov matrix used previously, and $\Sigma: p \times p$ is positive definite. The vector $\beta$ and $\Sigma$ are unknown.

3.6.2 Estimation

Based on the matrix normal distribution, the likelihood function for the model in (3.5) is

$$L(\beta, \Sigma) \propto |\Sigma|^{-\frac{1}{2} n} e^{-\frac{1}{2} tr(\Sigma^{-1}(X-\hat{A}\beta_1')(X-\hat{A}\beta_1')')},$$  \hspace{1cm} (3.6)

where $L(\beta, \Sigma)$ denotes the likelihood function with the parameters $\beta$ and $\Sigma$. Note that $A = \Sigma G_a$. To obtain the estimator of $\Sigma$ and $\beta$ through the likelihood function is not a trivial task, since $\Sigma$ appears both in the mean and the variance. However, we manage to derive the explicit MLEs for given $\omega$ via inequalities, as formulated in the next theorem.

**Theorem 3.1.** Let the model be given by (3.5) and suppose that $\omega$ in $A$ is known, with $A = \Sigma G_a = (\Sigma\omega, \Sigma^2\omega, \ldots, \Sigma^a\omega)$, and that $S = X(I - 1_n 1_n'n^{-1})X'$. Then, if $n > p$, the maximum likelihood estimators of $\Sigma$ and $A\beta$ are given by

$$\hat{A}\beta = \hat{A}(\hat{A}'S^{-1}\hat{A})^{-1}\hat{A}'S^{-1}X1_n1_n'n^{-1};$$

$$\hat{A} = (\frac{1}{n}S\omega, \frac{1}{n^2}S^2\omega, \ldots, \frac{1}{n^a}S^a\omega),$$

$$\hat{\Sigma} = \frac{1}{n}\{S+(I-\hat{A}(\hat{A}'S^{-1}\hat{A})^{-1}\hat{A}'S^{-1})X1_n1_n'n^{-1}X'(I-S^{-1}\hat{A}(\hat{A}'S^{-1}\hat{A})^{-1}\hat{A}').

**Proposition 3.1.** Assume that $\omega$ and $\mu_y$ are known and the given observations $X$ follow the model in (3.5). The prediction of $y$ is given by

$$\hat{y}' = \omega'\hat{\Sigma}^{-1}(X - \hat{A}\beta_1') + \mu_y,$$  \hspace{1cm} (3.7)

and $\hat{A}\beta$ is presented in Theorem 3.1.
The details of the proof of Theorem 3.1 are given in Paper II. Explicit MLEs of $\Sigma$ and $\beta$ will simplify computations. Observe that in the MLEs of $A$, i.e. $(\Sigma, \Sigma^2, \cdots, \Sigma^a)$, every $\Sigma$ is replaced with the sample variance, i.e. $S/n$. The MLE of $\Sigma$ differs from the sample variance, which leads to the estimator being biased. However, we have argued (for details see in Paper III) that the bias will not cause any problem in prediction. The estimation obtained through the two-step method will always be better than that obtained using PLS in that there will be a smaller mean squares error, which was shown in Paper III.

3.6.3 Data example

The data of one silage experiment consisting of 215 samples were used for illustrating the performance of the two-step method. The ethanol content in the silage was the response variable and the absorbance values of the spectra from the Fourier transform infrared (FTIR) analysis at 53 wave numbers were the predictors. The data set was randomly divided into two sets. One set consisting of 115 samples was used to perform the regression analysis. The other 100 samples were used for validation. The estimation and prediction results are shown in Figure 8. The two-step method gave better estimation and prediction than PLS when the number of components was small, i.e. less than eight in this case. However, both methods performed similarly when more components were included in the model. Fearn’s data (Fearn, 1983) and a simulation study were included in Paper III, where similar conclusions were made.

![Figure 8](image_url): The square root of the mean squared error of estimation (MSE) and the square root of the mean squared error of prediction (PSE) for PLS and the two-step method for the data from one selected silage experiment. The left figure shows the root of the MSE, while the right figure shows the root of the PSE.
3.7 The two-step method for group effects

Formulating PLS as a two-step method gives a great modelling flexibility, making it possible, for example, to combine the data from different studies. Due to the experimental conditions, data from different treatment groups are often collected, e.g. groups representing different genders, seasons, etc. In silage data, chemical compounds such as sucrose can naturally be high in one experiment, but relatively low in another experiment, due to the crop type used. An illustration of data with group effects is provided in Figure 9. The curves of the two groups in this figure show common peaks, which indicates that the $X$ are governed by the same underlying components. In addition, the curves from the two groups have two distinct levels, which indicates that the group effects on the mean of the two experiments are different.

Figure 9: Data with group effects. Each group is represented by one colour. The peaks of the curves indicate underlying effects. The two curves share the same peaks, but have different levels, which implies that the mean structure is the same, but has two distinct levels.

3.7.1 Model

Following the idea of the classical growth curve model, the mean of $X$ for each group is supposed to be linear with the underlying components. Thus, the first step model for the group effect turns into:

$$X = ABC + E,$$

(3.8)

with $X$ and $A$ having the same definition as in the model in (3.5), and with $B$: $q \times k$, $C$: $k \times n$, where $k$ is the number of groups. $A$ is still the within-individuals design matrix, which indicates that each group shares the same underlying structure. The matrix $C$ is the between-individuals design matrix, which keeps track of the observations in the groups. For example, if there are
three groups with three, five and four observations in each group, we may use

$$C = \begin{pmatrix}
1 & 1 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 1 & 1 & 1 & 1 & 1 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 1 & 1 & 1
\end{pmatrix},$$

and if \( k = 1 \) (i.e. one group), then \( C = 1_n' \). Moreover, \( E \sim N_{p,n}(0, \Sigma, I_n) \), and \( \Sigma: p \times p \) is supposed to be positive definite. The matrices \( B \) and \( \Sigma \) are unknown and should be estimated. The model in (3.5) is a special case of that in (3.8) when \( k = 1 \).

### 3.7.2 Estimation

The likelihood function for the model in (3.8) will be identical to the one for the model in (3.5) by replacing \( 1_n' \) with \( C \). The technique for obtaining MLEs in the model in (3.5) is also applicable to the estimation here. The estimation results are summarized in the next theorem, which is one of the main results of Paper III.

**Theorem 3.2.** Let the model be given by (3.8) and suppose that \( \omega \) in \( A \) is known, with \( A = \Sigma G_a = (\Sigma \omega, \Sigma^2 \omega, \ldots, \Sigma^a \omega) \), and that \( S = X(I - P_{C'})X' \), where \( P_{C'} = C'(CC')^{-1}C \). Then, if \( n > p \), the maximum likelihood estimators of \( \Sigma \) and \( AB \) are given by

$$\hat{AB} = \hat{A}(\hat{A}'S^{-1}\hat{A})^{-1}\hat{A}'S^{-1}XC'(CC')^{-1},$$

$$\hat{A} = (\frac{1}{n}S\omega, \frac{1}{n^2}S^2\omega, \ldots, \frac{1}{n^a}S^a\omega),$$

$$\hat{\Sigma} = \frac{1}{n}(S + (I - \hat{A}(\hat{A}'S^{-1}\hat{A})^{-1}\hat{A}'S^{-1})XP_{C'}X' \times (I - S^{-1}\hat{A}(\hat{A}'S^{-1}\hat{A})^{-1}\hat{A}')).$$

**Proposition 3.2.** Assume that \( \omega \) and \( \mu_y \) are known and the given observations \( X \) follow the model in (3.8). The prediction of \( y \) is

$$\hat{y}' = \omega'\hat{\Sigma}^{-1}(X - \hat{\mu}_x C) + \mu_y', \quad \hat{\mu}_x = \hat{AB}, \quad (3.9)$$

and \( \hat{AB} \) is given in Theorem 3.2.

Details of derivation of Theorem 3.2 are presented in Paper III. \( \hat{\Sigma} \) in Theorem 3.2 is also a biased estimator, but this will not cause any serious problem in prediction if the sample is large enough. A detailed discussion of this is given in Paper III. Theoretical comparisons of the two-step method and PLS become difficult in this case, since classical PLS does not take any group effect into account. Therefore, we will rely on some numerical results in the next subsection to obtain a general overview.
3.7.3 Data example

In order to illustrate the performance of the two-step method for grouped data, two experiments (referred to as Exp 1 and Exp 2) were selected. The ethanol content was the response variable and 53 wave numbers were the predictors. The samples were divided into two parts. One part consisted of 265 samples (67 from Exp 1 and 198 from Exp 2) which were used for building the model. The other part, consisting of 40 samples, 16 of which came from Exp 1, was used for testing the model performance.

The mean ethanol contents of the samples of Exp 1 and Exp 2 used for building the model were 2.3 g/L and 4.8 g/L, respectively. Part of the spectra has been displayed in Figure 5, which indicates that there was a group effect. The estimation and prediction results are shown in Figure 10 and 11. The two-step method gave both better estimation and prediction than PLS. When including more components in the model, the superiority of the two-step method over PLS diminished, but did not cease, as highlighted in Figure 10 and 11. A similar phenomenon was also observed in a comprehensive simulation study in Paper III.

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**Figure 10:** The square root of the mean squared error of estimation (MSE) for PLS and the two-step method for data from two selected silage experiments. The left figure shows the square root of the MSE for the models consisting of one to ten components, respectively, while the right figure shows the square root of the MSE for the models consisting of 11 to 20 components, respectively (i.e. each successive model consists of one more component than the previous one).

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3.8 The two-step method for nested group effects

As noted before, grouped data can have a rather complicated structure besides exhibiting different mean levels; for example such data can be linear in one group and non-linear in another group, etc., as illustrated by Potthoff and Roy’s data in Figure 2. In Paper IV, we focused our modelling on nested group effects, i.e. the mean structure of one group being nested within another.
Figure 11: *The square root of the mean squared error of prediction (PSE) for PLS and the two-step method for data from two selected silage experiments.* The left figure shows the square root of the PSE for the models consisting of one to ten components, respectively, while the right figure shows the square root of the PSE for the models consisting of 11 to 20 components, respectively (i.e. each successive model consists of one more component than the previous one).

group. For example, part of the silage data concerns samples which have been subjected to special treatments. Consequently, there are relatively high levels of some compounds, e.g. ammonia, appearing in some experiments, whereas the same compounds are absent or occur in very low amounts in other experiments. The character of such data is illustrated in Figure 12. Each curve represents a group. The two curves have one peak in common, which indicates that there are the same underlying components. It is important to notice that one curve has one peak which is not exhibited by the other one. This peak corresponds to some underlying components which only play a role in one group. The matrix $A_1$ is used for modelling the common underlying components and $A_2$ is used for modelling the unique components which only matter for one of the groups (Group 2). The mean for Group 1 equals $\mu_1 = \beta_{10} + A_1 \beta_{11}$ and that for Group 2 is given by $\mu_2 = \beta_{20} + A_1 \beta_{21} + A_2 \beta_{22}$. Therefore, we can use the knowledge about the extended growth curve model to put the two groups together and create one general model, which will be presented in next subsection. The advantage of the new general model is that information from all the groups can be used to estimate $\Sigma$.

### 3.8.1 Model

We now present the set-up of the model for nested effects in three groups, i.e. $k = 3$. With $n$ observations, $n_i$ from group $i$ and $\sum_{i=1}^{3} n_i = n$, the model becomes:

$$X = A_1 B_1 C_1 + A_2 B_2 C_2 + A_3 B_3 C_3 + E, \quad E \sim N_{p,n}(0, \Sigma, I_n),$$

(3.10)
Figure 12: Data with nested group effects. Each group is represented by one colour. The peaks of the curves indicate underlying effects. There is an extra peak in one curve, which indicates that the mean structure of one group is more complex than that of the other one.

where,

\[ A_1 = (\Sigma \omega : \Sigma^2 \omega : \cdots : \Sigma^{a_1} \omega), \]
\[ A_2 = (\Sigma^{a_1+1} \omega : \Sigma^{a_1+2} \omega : \cdots : \Sigma^{a_1+a_2} \omega), \]
\[ A_3 = (\Sigma^{a_1+a_2+1} \omega : \Sigma^{a_1+a_2+2} \omega : \cdots : \Sigma^{a_1+a_2+a_3} \omega), \]
\[ C_1 = \begin{pmatrix} 1 & 1 & \cdots & 1 & 0 & 0 & \cdots & 0 & 0 & 0 & \cdots & 0 \\ 0 & 0 & \cdots & 0 & 1 & 1 & \cdots & 1 & 0 & 0 & \cdots & 0 \\ 0 & 0 & \cdots & 0 & 0 & 0 & \cdots & 0 & 1 & 1 & \cdots & 1 \end{pmatrix} , \]
\[ C_2 = \begin{pmatrix} 0 & 0 & \cdots & 0 & 1 & 1 & \cdots & 1 & 0 & 0 & \cdots & 0 \\ 0 & 0 & \cdots & 0 & 0 & 0 & \cdots & 0 & 1 & 1 & \cdots & 1 \end{pmatrix} , \]
\[ C_3 = \begin{pmatrix} 0 & 0 & \cdots & 0 & 0 & 0 & \cdots & 0 & 1 & 1 & \cdots & 1 \end{pmatrix} . \]

Moreover, \( \Sigma \) is supposed to be positive definite. The matrix \( B_i, i = 1, 2, 3, \) and \( \Sigma \) are the parameters which are to be estimated. Note that \( A_i \) is a function of \( \Sigma \). It is worth noting that the design matrices for Group 1-3 are \( A_1, (A_1 : A_2) \) and \( (A_1 : A_2 : A_3) \), respectively. For the form of a general model for nested group effects, we refer to Paper IV.

3.8.2 Estimation

The likelihood function for the model in (3.10) satisfies

\[
L(B_1, B_2, B_3, \Sigma) \propto |\Sigma|^{-\frac{1}{2}} n \text{etr} \{-\frac{1}{2} \Sigma^{-1} (X - \sum_{i=1}^{3} A_i B_i C_i)' \}, \tag{3.11}
\]
Theorem 3.3. Let the model be given by (3.10) and

\[
\begin{align*}
T_1 &= I - P_{A_1,\Sigma}, \quad T_2 = I - P_{T_1A_2,\Sigma}, \quad T_3 = I - P_{T_2T_1A_3,\Sigma}, \\
S_1 &= X(I - P_{C'_1})X', \quad S_2 = S_1 + T_1XP_{C'_1}(I - P_{C'_3})P_{C'_1}X'T_1', \\
S_3 &= S_2 + T_2T_1XP_{C'_1}P_{C'_2}(I - P_{C'_3})P_{C'_2}P_{C'_1}X'T_1'T_2', \\
Q &= A_1(\Sigma^{-1}A_1)^{-1}A_1' + T_1S_2/n, \\
M &= A_1(\Sigma^{-1}A_1)^{-1}A_1' + T_1A_2(\Sigma^{-1}T_1A_2)^{-1}A_2'T_1' - A_1(\Sigma^{-1}A_1)^{-1}A_1\Sigma^{-1}T_1A_2(\Sigma^{-1}T_1A_2)^{-1}A_2T_1' + T_1T_2S_3/n.
\end{align*}
\]

Suppose that \( \omega \) in \( A_i, i = 1, 2, 3, \) is known. Then, if \( n > p \), the maximum likelihood estimators of \( B_1, B_2, B_3 \) and \( \Sigma \) are given by

\[
\begin{align*}
\hat{B}_3 &= (\hat{A}_3\hat{T}_1\hat{T}_2\hat{S}_3^{-1}\hat{T}_2\hat{T}_1\hat{A}_3)^{-1}\hat{A}_3\hat{T}_1\hat{T}_2\hat{S}_3^{-1}\hat{T}_2\hat{T}_1XC_3'(C_3C_3')^{-1} \\
&\quad + (\hat{A}_3\hat{T}_1\hat{T}_2)^oZ_{31} + \hat{A}_3\hat{T}_1\hat{T}_2Z_{32}C_3'^o \\
\hat{B}_2 &= (\hat{A}_2\hat{T}_1\hat{S}_2^{-1}\hat{T}_1\hat{A}_2)^{-1}\hat{A}_2\hat{T}_1\hat{S}_2^{-1}(\hat{T}_1XP_{C'_2} - \hat{T}_1A_3\hat{B}_3C_3)C_2'(C_2C_2'^o)^{-1} \\
&\quad + (\hat{A}_2\hat{T}_1)^oZ_{21} + \hat{A}_2\hat{T}_1Z_{22}C_2' \\
\hat{B}_1 &= (\hat{A}_1\hat{S}_1^{-1}\hat{A}_1)^{-1}\hat{A}_1\hat{S}_1^{-1}(X - \hat{A}_2\hat{B}_2\hat{C}_2 - \hat{A}_3\hat{B}_3\hat{C}_3)C_1'(C_1C_1')^{-1} \\
&\quad + \hat{A}_1^oZ_{11} + \hat{A}_1^oZ_{12}C_1' \\
n\hat{\Sigma} &= \hat{S}_3 + \hat{T}_3\hat{T}_2\hat{T}_1XP_{C'_3}X'T_1'T_2'T_3',
\end{align*}
\]

where \( Z_{ij}, i = 1, 2, 3, j = 1, 2, \) are arbitrary matrices, \( C^o \) denotes any matrix.
satisfying $\zeta(C^\omega) = \zeta(C)^\perp$, and $\perp$ denotes the orthogonal complement,

\[
\begin{align*}
\hat{A}_1 &= \left(\frac{1}{n}S_1\omega : \frac{1}{n^2}S_1^2\omega : \cdots : \frac{1}{n^{a_1}}S_1^{a_1}\omega\right), \\
\hat{T}_1 &= I - \hat{A}_1(\hat{A}_1S_1^{-1}\hat{A}_1)^{-1}\hat{A}_1S_1^{-1}, \\
\hat{S}_2 &= S_1 + \hat{T}_1X P_{C_1}'(I - P_{C_2}')P_{C_1}'X\hat{T}_1, \\
\hat{Q} &= \hat{A}_1(\hat{A}_1S_1^{-1}\hat{A}_1)^{-1}\hat{A}_1' + \hat{T}_1\hat{S}_2/n, \\
\hat{A}_2 &= (\hat{Q}S_1^{a_1}\omega : \hat{Q}S_1^{a_1}\omega : \cdots : \hat{Q}S_1^{a_1}\omega)/n^{a_1}, \\
\hat{T}_2 &= I - \hat{T}_1\hat{A}_2(\hat{A}_2'\hat{T}_1'\hat{S}_2'^{-1}\hat{T}_1'\hat{A}_2')^{-1}\hat{A}_2'\hat{T}_1'^{-1}, \\
\hat{S}_3 &= \hat{S}_2 + \hat{T}_2\hat{T}_1'X P_{C_1}'P_{C_2}'(I - P_{C_3}')P_{C_1}'X\hat{T}_1'\hat{T}_2', \\
\hat{M} &= \hat{A}_1(\hat{A}_1S_1^{-1}\hat{A}_1)^{-1}\hat{A}_1' + \hat{T}_1\hat{A}_2(\hat{A}_2'\hat{T}_1'\hat{S}_2'^{-1}\hat{T}_1'\hat{A}_2')^{-1}\hat{A}_2'\hat{T}_1' \\
&\quad - \hat{A}_1(\hat{A}_1S_1^{-1}\hat{A}_1)^{-1}\hat{A}_1'\hat{T}_1^{-1}\hat{A}_2(\hat{A}_2'\hat{T}_1'\hat{S}_2'^{-1}\hat{T}_1'\hat{A}_2')^{-1}\hat{A}_2'\hat{T}_1', \\
\hat{A}_3 &= \left(\hat{M}'\hat{Q}^{a_2}S_1^{a_1}\omega : \hat{M}'\hat{Q}^{a_2}S_1^{a_1}\omega : \cdots : \hat{M}'\hat{Q}^{a_2}S_1^{a_1}\omega\right)/n^{a_1}. 
\end{align*}
\]

**Proposition 3.3.** Assume that $\omega$ and $\mu_y$ are known and the given observations $X$ follow the model in (3.10) with $k = 3$. The prediction of $y$ equals:

\[
y = \omega'\hat{\Sigma}^{-1}(X - \hat{A}_1\hat{B}_1C_1 - \hat{A}_2\hat{B}_2C_2 - \hat{A}_3\hat{B}_3C_3) + \mu_y,
\]

and $\hat{A}_i\hat{B}_i$ is given in Theorem 3.3.

We can observe that the MLEs of $B_i$ are given in ambiguous forms which depends on arbitrary matrices. However, it can be shown that $\hat{A}_1\hat{B}_1C_1 + \hat{A}_2\hat{B}_2C_2 + \hat{A}_3\hat{B}_3C_3$ is unique, which indicates that the estimations of the mean and future prediction are unique.

### 3.8.3 Data example

Two experiments, which will be referred to as Exp 3 and Exp 4, were selected for illustrating the performance of the method. The corresponding spectra for these two experiments have been shown in Figure 6. It is clear that there is an underlying structure in one experiment (Exp 4) which does not occur in the other one (Exp 3). The lactate content was the response variable and 53 wave numbers were the predictors. Exp 3 was divided into two parts, one of which had 80 samples for training, i.e. building the model, while the other part had 20 samples for testing, i.e. checking the model performance. Exp 4 was also divided into two parts, a training part consisting of 50 samples and a testing part consisting of 21 samples.

First the data were analyzed “marginally”; i.e. one model was built by only using the 80 training samples in Exp 3 and then tested using the remaining
20 testing samples in Exp 3. The same procedure was applied to Exp 4. The prediction results of the two-step method are shown in Figure 13. The prediction error for Exp 3 becomes stable after including just one component in the model, while the prediction error for Exp 4 becomes stable after including at least three components. Thus the “marginal” analysis suggests that these two experiments require different numbers of components.

![Figure 13: The square root of the mean squared error of prediction (PSE) for the two-step method for data from two selected silage experiments. The left figure is for Exp 3 and the right figure for Exp 4.](image)

Furthermore, we examined the model with a group effect by putting the two experiments together, i.e. building the regression model using the training samples from Exp 3 and Exp 4. As shown in Figure 14, after including five components in the model, i.e. $A = \Sigma G_5$ in the model in (3.8), the prediction error obtained when using the “combined” data is better than that obtained when using the experiments separately.

Finally, the model in (3.10) with the nested mean structure was used by specifying some additional components in Exp 4. Correspondingly, the model in (3.10) became $X = A_1 B_1 C_1 + A_2 B_2 C_2 + E$, with $A_1 = \Sigma G_{a_1}$ and $A_2 = \Sigma G_{a_2}$. One additional component was assigned for Exp 4, i.e. $a_2 = 1$. Based on the prediction error (as shown in Figure 15), very little improvement is gained by using the model in (3.10) compared to using the model in (3.8) when only one additional component is included in Exp 4.
Figure 14: The square root of the mean squared error of prediction (PSE) for the two-step method for data from two selected silage experiments using “marginal” or “combined” analysis.

Figure 15: The square root of the mean squared error of prediction (PSE) for the two-step method for data from two selected silage experiments using the group effect model in (3.8) and the nested effect model in (3.10).
4 Conclusions, discussion and future work

4.1 Contributions

The main contribution of the thesis is that PLS has been put into a classical multivariate regression model together with a two-step prediction approach. Consequently, in comparison with PLS, the two-step method is non-algorithmic and possesses greater modelling flexibility. After the connections were established, the two-step method was extended to model data with more complex structures, i.e. with a group effect and with a nested mean structure. Under the two-step method and its extensions, explicit maximum likelihood estimators have been derived.

The contributions of each paper are summarized in detail as follows:

Paper I:
- The robustness of the popular shrinkage methods was investigated through their applications on real silage data with a rather complex structure, i.e. multi-responses and group effects.
- Two simulation procedures were included in the comparisons. One checked the prediction for the randomly selected samples, while the other one focused on the prediction for samples from a different experiment.

Paper II:
- A two-step method for linear prediction, especially for collinear data, is proposed.
- PLS is linked to the two-step method via a Krylov design matrix; i.e. in the first step the explanatory variables are summarized via a multilinear model with a Krylov structured matrix.
- Given the covariance $\omega$, explicit maximum likelihood estimators for the mean and dispersion were derived.

Paper III:
- The connections between PLS and the two-step method give a new way of viewing PLS. PLS can be considered as a method of approximating $\Sigma^{-1}$ using the Cayley-Hamilton theorem.
- The two-step method was extended to combine data from different studies, by including a between-individuals design matrix.
- Explicit maximum likelihood estimators for the group mean and the dispersion matrix of the explanatory variables were derived.
- The properties of the estimators, e.g. the bias, were discussed. It was shown that for the within-sample prediction, the mean squared error of the two-step method is always smaller than that of PLS.
• Numerical illustration using a real data set and simulated data were included to assess the performance of the two-step method in various cases, in comparison with that of PLS and other regularization methods. The numerical results indicate that the two-step method outperforms the other methods in the case of grouped data.

Paper IV:

• The two-step method was extended to model grouped data which, besides having collinear explanatory variables, possesses a nested mean structure. In the first step, the explanatory variables were modelled in a multilinear model with a structure similar to the extended growth curve structure, but with Krylov design matrices which depend on $\Sigma$.

• Surprisingly, under such a general multilinear model structure, explicit maximum likelihood estimators for the mean and dispersion matrices were derived.

4.2 Discussion

PLS inspired us to start the two-step method with $x = \omega \gamma + \varepsilon$, which assumes that the predictors are proportional to the covariance. Despite the fact that such a model works according to the theoretical and numerical results, the reason why such an assumption works is still mysterious. The connection between PLS and the two-step method also led us to choose $\Sigma G_a$ as the design matrix. However, why does this particular choice work so well? It is unclear what information is included in the Krylov space. The iterative algorithm of PLS indicates that the previously generated weights $\omega_i$ play a more important role than the subsequently generated ones. Therefore, the corresponding elements $\Sigma^{i-1} \omega$ in the Krylov space should be more important than the subsequent ones, which is not obvious. According to a numerical pilot study, it seems that a subsequently generated component is not always smaller than the previously generated ones. In our two-step method, the MLEs have played the role of being a weight factor, i.e. weighting each element in the Krylov space to fit the mean.

Compared with the envelopes of Cook et al. (2010, 2013), the two-step method is more specific. The two-step method gives explicit MLEs using a semi-population set-up where the covariance between the response and the predictor and the mean of the response are known, while the MLEs for all the parameters in envelopes are determined numerically. Using the conditional distribution, it is possible to derive the covariance and the mean of the response in the two-step method through the maximum likelihood approach. We foresee that the estimators will depend on numerical optimizations, which may affect the finding of explicit MLEs and create a burden of computation. For example, the computation will become very slow in the optimization of the likelihood using the envelope method when $p$ is large, whereas the explicit MLEs in the two-step method do not suffer from such a problem. Therefore,
it is not obvious in the two-step method which estimator is better, i.e. the explicit MLEs using semi-population model or MLEs for all the parameters.

4.3 Future work

Stopping rules, i.e. rules concerning how many terms should be included in $A = \Sigma G_a$, represent one of the most interesting questions to be investigated in the future. The most commonly applied approach is to use cross-validation, which is very computationally demanding. If PLS stops, the Krylov space, which is included in the design matrix of $A$, turns out to be an invariant space, and the space of a lower or an equal dimension compared to the original one. Therefore, a future topic of research would be the definition of an optimum stopping rule for the selection of an appropriate model based on the Krylov space.

As mentioned earlier, the two-step method has a great modelling flexibility. We have developed models for grouped data. Another extension would be to include additional covariates. For example, we are still interested in building a prediction model for the concentrations of some chemical compound, $y$, from the spectra $X$. Then the model in the first step would be $X = A_1 B_1 C_1 + E$, where $A_1 = \Sigma G_a$ and $C_1$ contains group information, e.g. three groups, each of which has three, five, four observations. In addition, the temperature may also influence the spectra. Then, the model could be extended as $X = A_1 B_1 C_1 + B_2 C_2 + E$, where $C_2$ contains the temperature information. Note that the temperature can vary in the group, i.e.

$$C_2 = \begin{pmatrix} 15 & 10 & 10 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 20 & 20 & 20 & 15 & 15 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 10 & 16 & 16 & 16 \end{pmatrix}.$$

The estimators in such a model could be obtained.

Partial least squares have been found useful in a wide range of statistical fields, e.g. discriminant analysis, logistic regression, etc. A natural future area of research would be an extension of the two-step method for implementation in these fields, which would lead to new applications.
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