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Simultaneous Dimensionality Reduction: A Data Efficient Approach for Multimodal Representations Learning

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Simultaneous Dimensionality Reduction: A Data Efficient Approach for Multimodal Representations Learning

By

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Abstract

Simultaneous Dimensionality Reduction: A Data Efficient Approach for Multimodal Representations Learning
By Eslam Abdelaleem

Current experiments frequently produce high-dimensional, multimodal datasets—such as those combining neural activity and animal behavior or gene expression and phenotypic profiling—with the goal of extracting useful correlations between the modalities. Often, the first step in analyzing such datasets is dimensionality reduction. We explore two primary classes of approaches to dimensionality reduction (DR): Independent Dimensionality Reduction (IDR) and Simultaneous Dimensionality Reduction (SDR). In IDR methods, of which Principal Components Analysis is a paradigmatic example, each modality is compressed independently, striving to retain as much variation within each modality as possible. In contrast, in SDR, one simultaneously compresses the modalities to maximize the covariation between the reduced descriptions while paying less attention to how much individual variation is preserved. Paradigmatic examples include Partial Least Squares and Canonical Correlations Analysis. Even though these DR methods are a staple of statistics, their relative accuracy and data set size requirements are poorly understood. We introduce a generative linear model to synthesize multimodal data with known variance and covariance structures to examine these questions. We assess the accuracy of the reconstruction of the covariance structures as a function of the number of samples, signal-to-noise ratio, and the number of varying and covarying signals in the data. Using numerical experiments, we demonstrate that linear SDR methods consistently outperform linear IDR methods and yield higher-quality, more succinct reduced-dimensional representations with smaller datasets. Remarkably, regularized CCA can identify low-dimensional weak covarying structures even when the number of samples is much smaller than the dimensionality of the data, which is a regime challenging for all dimensionality reduction methods. Our work corroborates and explains previous observations in the literature that SDR can be more effective in detecting covariation patterns in data. These findings suggest that SDR should be preferred to IDR in real-world data analysis when detecting covariation is more important than preserving variation.

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Chapter 1

Introduction

Many modern experiments across various fields generate massive multimodal data sets. For instance, in neuroscience, it is common to record the activity of a large number of neurons while simultaneously recording the resulting animal behavior [44, 43, 46, 30]. Other examples include measuring gene expressions of thousands of cells and their corresponding phenotypic profiles, or integrating gene expression data from different experimental platforms, such as RNA-Seq and microarray data [10, 51, 45, 27, 31]. In economics, important variables such as inflation are often measured using combinations of macroeconomic indicators as well as indicators belonging to different economic sectors [20, 2, 15, 40]. In all of these examples, an important goal is to estimate statistical correlations among the different modalities.

Analyses usually begin with dimensionality reduction (DR) into a smaller and more interpretable representation of the data. We distinguish two types of DR: *independent* (IDR) and *simultaneous* (SDR) [34]. In the former, each modality is reduced independently, while aiming to preserve its variation, which we call *self* signal. In the latter, the modalities are compressed simultaneously, while maximizing the covariation (or the *shared* signal) between the reduced descriptions and paying less attention to preserving the individual variation. It is not clear if IDR techniques, such as the

Principal Components Analysis (PCA) [25], are well-suited for extracting shared signals since they may overlook features of the data that happen to be of low variance, but of high covariance [11]. In particular, poorly sampled weak shared signals, common in high-dimensional datasets, can exacerbate this issue. SDR techniques, such as Partial Least Squares (PLS) [50] and Canonical Correlations Analysis (CCA) [26], are sometimes mentioned as more accurate in detecting weak shared signal [9, 21, 36]. However, the relative accuracy and data set size requirements for detecting the shared signals remain poorly understood for both classes of methods.

In this study, we aim to assess the strengths and limitations of linear IDR, represented by PCA, and linear SDR, exemplified by PLS and CCA, in detecting weak shared signals. For this, we introduce a generative linear model that captures key features of relevant examples, including noise, the self signal, and the shared signal components. Using this model, we analyze the performance of the methods in different conditions. Our goal is to assess how well these techniques can (i) extract the relevant shared signal and (ii) identify the dimensionality of the shared and the self signals from noisy, undersampled data. We investigate how the signal-to-noise ratios, the dimensionality of the reduced variables, and the method of computing correlations combine with the sample size to determine the quality of the DR. We propose best practices for achieving high-quality reduced representations with small sample sizes using these linear methods.

Chapter 2

Model

2.1 Relations to Previous Work

The extraction of signals from large-dimensional data sets is a challenging task when the number of observations is comparable to or smaller than the dimensionality of the data. The undersampling problem introduces spurious correlations that may appear as signals, but are, in fact, just statistical fluctuations. This poses a challenge for DR techniques, as they may retain unnecessary dimensions or identify noise dimensions as true signals.

Here, we focus exclusively on linear DR methods. For these, the Marchenko-Pastur (MP) distribution of eigenvalues of the covariance matrix of pure noise derived using the Random Matrix Theory methods [33] has been used to introduce a cutoff between noise and true signal in real datasets. However, recent work [13] has shown that, when observations are a linear combination of uncorrelated noise and latent low-dimensional self signals, then the self signals alter the distribution of eigenvalues of the sampling noise, questioning the validity of this naive approach.

Moving beyond a single modality, [4] calculated the singular value spectrum of cross-correlations between two nominally uncorrelated random signals. However, it remains unknown whether the linear mixing of self signals and shared signals affects the singular value spectra of noise, and how all of these components combine to limit the ability to detect shared signals between two modalities from data sets of realistic sizes. Filling in this gap using numerical simulations is the main goal of this paper, and analytical treatment of this problem will be left for the future.

The linear model and linear DR approaches studied here do not capture the full complexity of real-world data sets and state-of-the-art algorithms. However, if sampling issues and self signals limit the ability of linear DR methods to extract shared signals, it would be surprising for nonlinear methods to succeed in similar scaling regimes on real data. Thus extending the previous work to explicitly study the effects of linear mixtures of self signals, shared signals, and noise on limitations of DR methods is likely to endow us with intuition that is useful in more complex scenarios routinely encountered in different domains of science.

Examples of scenarios with shared and self signals include inference of dynamics of a system through a latent space [12, 8], where shared signals correspond to latent factors that are relevant for predicting the future of the system from its past, while self signals correspond to nonpredictive variation [3]. In economics, shared and self signals correspond to diverse macroeconomic indicators that are grouped into correlated distinct categories in structural factor models [14, 20, 40, 2]. In neuroscience, shared signals can correspond to the latent space, by which neural activity affects behavior, while self signals encode neural activity that does not manifest in behavior and behavior that is not controlled by the part of the brain being recorded from [42, 44, 35, 41, 37, 46, 30].

Interestingly, in the context of the neural control of behavior, it was noticed that SDR reconstructs the shared neuro-behavioral latent space more efficiently and using a smaller number of samples than IDR [41]. Similar observations have been made in more general statistical contexts [9, 21, 36], though the agreement is not

uniform [17, 18, 19]. Because of this, most practical recommendations for detecting shared signals are heuristic [22], with widely acknowledged, but poorly understood limitations and possible resolutions [29]. Our goal is to ground such rules in numerical simulations and scaling arguments.

2.2 Linear Model with Self and Shared Signals

We consider a linear model with noise, $m_{\rm self,X}$, $m_{\rm self,Y}$ self signals that are relevant to each modality independently, as well as $m_{\rm shared}$ shared signals that capture the interrelationships between modalities. It results in T observations of two high-dimensional standardized observables, X and Y:

$$\begin{bmatrix} \tilde{X} \in \mathbb{R}^{N_X} \end{bmatrix} = \underbrace{R_X}_{\text{Independent white noise}} + \underbrace{U_X V_X}_{\text{Self-Signal for X}} + \underbrace{PQ_X}_{\text{Shared-Signal}},$$

$$\begin{bmatrix} \tilde{Y} \in \mathbb{R}^{N_Y} \end{bmatrix} = \underbrace{R_Y}_{\text{Independent white noise}} + \underbrace{U_Y V_Y}_{\text{Self-Signal for Y}} + \underbrace{PQ_Y}_{\text{Shared-Signal}},$$

$$X = \tilde{X} / \sigma_{\tilde{X}}, Y = \tilde{Y} / \sigma_{\tilde{Y}}. \tag{2.2}$$

The observations of X and Y are linear combinations of the following: (a) Independent white noise components R_X and R_Y with variances $\sigma_{R_X}^2$ and $\sigma_{R_Y}^2$. (b) Self-signal components U_X and U_Y residing in lower-dimensional subspaces $\mathbb{R}^{m_{\text{self},Y}}$ and $\mathbb{R}^{m_{\text{self},Y}}$ with variances $\sigma_{U_X}^2$ and $\sigma_{U_Y}^2$. (c) Shared-signal components P in a shared lower-dimensional subspace $\mathbb{R}^{m_{\text{shared}}}$ with variance σ_P^2 . These components are projected into their respective high-dimensional spaces \mathbb{R}^{N_X} and \mathbb{R}^{N_Y} using fixed quenched projection matrices V_X , V_Y , Q_X , and Q_Y with specified variances $\sigma_{V_X}^2$, $\sigma_{V_Y}^2$, $\sigma_{Q_X}^2$, and $\sigma_{Q_Y}^2$, all respectively. Entries in these matrices are drawn from a Gaussian distribution with a zero mean and the corresponding variances. Further, division by $\sigma_{\tilde{X}}$ and $\sigma_{\tilde{Y}}$ standardizes each column of the data matrices by their empirical standard deviations. The total variance in the matrix \tilde{X} can be calculated as the sum of the variances of

its individual components: $\sigma_{\tilde{X}}^2 = \sigma_{R_X}^2 + m_{\text{self},X} \times \sigma_{U_X}^2 \sigma_{V_X}^2 + m_{\text{shared}} \times \sigma_P^2 \sigma_{Q_X}^2$. A similar calculation can be done for the total variance in \tilde{Y} .

We define self and shared signal-to-noise ratios $\gamma_{\text{self},X/Y}$, $\gamma_{\text{shared},X/Y}$ as the relative strength of signals compared to background noise per component in each modality. These definitions allow us to examine how easily self or shared signals in each dimension can be distinguished from the noise.

$$\gamma_{\text{self},X/Y} = \frac{\sigma_{U_{X/Y}}^2 \sigma_{V_{X/Y}}^2}{\sigma_{R_{X/Y}}^2}, \quad \gamma_{\text{shared},X/Y} = \frac{\sigma_P^2 \sigma_{Q_{X/Y}}^2}{\sigma_{R_{X/Y}}^2}$$
(2.3)

Our main goal is to evaluate the ability of linear SDR and IDR methods to reconstruct the shared signal P between X and Y while overlooking the effects of the self signals U_X, U_Y on the statistics of the shared ones.

Chapter 3

Methods

We apply DR techniques to X and Y to obtain their reduced dimensional forms Z_X and Z_Y , respectively. Z_X, Z_Y are of sizes that can range from $T \times 1$ to $T \times N_X$ and $T \times N_Y$, respectively. As an IDR method, we use PCA [25]. As SDR methods, we apply PLS [50] and CCA [26, 47, 52], including both normal and regularized versions of the latter. Each of these methods focuses on specific parts of the overall covariance matrix

$$C_{X,Y} = \begin{bmatrix} C_{XX} & C_{XY} \\ C_{YX} & C_{YY} \end{bmatrix} = \begin{bmatrix} \frac{1}{T}X^{\top}X & \frac{1}{T}X^{\top}Y \\ \frac{1}{T}Y^{\top}X & \frac{1}{T}Y^{\top}Y \end{bmatrix}.$$
(3.1)

PCA aims to identify the most significant features that explain the majority of the variance in C_{XX} and C_{YY} , independently. PLS, on the other hand, focuses on singular values and vectors that explain the covariance component C_{XY} . Along the same lines, CCA aims to find linear combinations of X and Y that are responsible for the correlation $(C_{XY}/\sqrt{C_{XX}C_{YY}})$ between X and Y. A detailed description of each method is in the next section 3.1.

For every numerical experiment, we generate training and test data sets $(X_{\text{train}}, Y_{\text{train}})$ and $(X_{\text{test}}, Y_{\text{test}})$ according to Eqs. (2.1-2.2). We apply PCA, PLS, CCA, and regularized CCA (rCCA) to the training to obtain the singular directions $W_{X_{\text{train}}}$ and

 $W_{Y_{\text{train}}}$ for each method (detailed in section 3.1). We then evaluate the projections of the test data on these singular directions

$$Z_{X_{\text{test}}} = X_{\text{test}} W_{X_{\text{train}}},$$

$$Z_{Y_{\text{test}}} = Y_{\text{test}} W_{Y_{\text{train}}}.$$
(3.2)

Finally, we evaluate the reconstructed correlations metric \mathcal{RC}' , which measures how well these singular directions recover the shared signals in the data, corrected by the expected positive bias due to the sampling noise, as detailed in section 3.2. $\mathcal{RC}' = 0$ corresponds to no overlap between the true and the recovered shared directions, and $\mathcal{RC}' = 1$ corresponds to perfect recovery.

3.1 Linear Dimensionality Reduction Methods

3.1.1 PCA

PCA is a widely used linear IDR method that aims to find the orthogonal principal directions, such that a few of them explain the largest possible fraction of the variance within the data. PCA decomposes the covariance matrix of the data matrix X, $C_{XX} = \frac{1}{T}X^{\top}X$, into its eigenvectors and eigenvalues through singular value decomposition (SVD). The SVD yields orthogonal directions, represented by the vectors $w_X^{(i)}$, that capture the most significant variability in the data. In most numerical implementations [38], these directions are obtained consecutively, one by one, such that the dot product between any two directions is zero $w_X^{(i)} \cdot w_X^{(j)} = \delta_{ij}$. The eigenvectors $w_X^{(i)}$ are obtained as the best solution to the optimization problem:

$$w_X^{*(i)} = \underset{w_X^{(i)}}{\arg\max} \frac{w_X^{(i)^{\top}} X^{(i)^{\top}} X^{(i)} w_X^{(i)}}{w_X^{(i)^{\top}} w_X^{(i)}}.$$
 (3.3)

Here $X^{(i)}$ is the *i*th deflated matrix where $X^{(1)}$ is the original matrix, and for every subsequent i+1, the matrix is deflated by subtracting the projection of X on the obtained weights: $X^{(i+1)} = X - \sum_{s=1}^{i} X w_{(s)} w_{(s)}^{\top}$. The eigenvectors are sorted in decreasing order according to their corresponding eigenvalues, and the first k eigenvectors $w_X^{(i=1:k)}$ are selected to form the projection matrix W_X . The obtained vectors determine the size of the reduced form Z_X , where $|Z_X| = k$ is the number of vectors retained from the decomposition of X. The vectors $w_X^{(i)}$ are then stacked together to form the projection matrix W_X . The low-dimensional representation Z_X is then obtained by multiplying the original data matrix X with this projection matrix, resulting in the reduced data matrix $Z_X = XW_X$. Similar treatment is done for Y in order to obtain $Z_Y = YW_Y$

One of the main advantages of PCA is its simplicity and efficiency. However, one of the drawbacks of this method is that it performs DR for X and Y independently, and one then searches for relations between Z_X and Z_Y by regressing one on the other. Thus obtained low-dimensional descriptions may capture variance but not the covariance between the two datasets.

3.1.2 PLS

PLS, or Partial Least Squares, performs SDR by finding the shared signals that explain the maximum covariance between two sets of data [50]. PLS performs the SVD of the covariance matrix $C_{XY} = \frac{1}{T}X^{\top}Y$ (or equivalently $C_{YX} = \frac{1}{T}Y^{\top}X$). The left and right singular vectors $(w_X^{*(i)}, w_Y^{*(i)})$ are obtained consecutively pair by pair such that $w_X^{(i)} \cdot w_Y^{(j)} = \delta_{ij}$. They are solutions of the optimization problem:

$$(w_X^{*(i)}, w_Y^{*(i)}) = \underset{w_X^{(i)}, w_Y^{(i)}}{\operatorname{arg max}} \frac{w_X^{(i)^{\top}} X^{(i)^{\top}} Y^{(i)} w_Y^{(i)}}{\sqrt{(w_Y^{(i)^{\top}} w_Y^{(i)})(w_Y^{(i)^{\top}} w_Y^{(i)})}}$$
(3.4)

The matrices $X^{(i)}, Y^{(i)}$ are deflated in a similar manner to PCA 3.1.1. The singular vectors are sorted in the decreasing order of their corresponding singular values, and the first k vectors are selected to form the projection matrices (W_X, W_Y) . The obtained vectors determine the size of the reduced form (Z_X, Z_Y) , where $|Z_X| = |Z_Y| = k$ is the number of vectors retained. The vectors $(w_X^{(i)}, w_Y^{(i)})$ are then stacked together to form the projection matrices (W_X, W_Y) respectively. The low-dimensional representations (Z_X, Z_Y) are obtained by projecting the original data matrices (X, Y) onto these projection matrices: $Z_X = XW_X$, and $Z_Y = YW_Y$.

In summary, PLS performs simultaneous reduction on both datasets, maximizing the covariance between the reduced representations Z_X and Z_Y . This property makes PLS a powerful tool for studying the relationships between two datasets and identifying the underlying factors that explain their joint variability.

3.1.3 CCA

CCA is another SDR method, which aims to find the directions that explain the maximum correlation between two datasets [26]. However, unlike PLS, CCA obtains the shared signals by performing SVD on the correlation matrix $\frac{C_{XY}}{\sqrt{C_{XX}}\sqrt{C_{YY}}}$. The singular vectors $(w_X^{*(i)}, w_Y^{*(i)})$ are obtained consecutively pair by pair such that $w_X^{(i)} \cdot w_Y^{(j)} = \delta_{ij}$. CCA enforces the orthogonality of $w_X^{(i)}, w_Y^{(i)}$ independently as well, such that $w_X^{(i)} \cdot w_X^{(j)} = w_Y^{(i)} \cdot w_Y^{(j)} = \delta_{ij}$. The singular vectors are obtained by solving the optimization problem:

$$(w_X^{*(i)}, w_Y^{*(i)}) = \underset{w_X^{(i)}, w_Y^{(i)}}{\operatorname{arg\,max}} \frac{w_X^{(i)^{\top}} X^{(i)^{\top}} Y^{(i)} w_Y^{(i)}}{\sqrt{(w_X^{(i)^{\top}} X^{(i)^{\top}} X^{(i)} w_X^{(i)})(w_Y^{(i)^{\top}} Y^{(i)^{\top}} Y^{(i)} w_Y^{(i)})}}.$$
 (3.5)

Like in PLS 3.1.2, the matrices $X^{(i)}$, $Y^{(i)}$ are deflated in a similar manner. In addition, the first k singular vectors $(w_X^{*(i)}, w_Y^{*(i)})$ are stacked together to form the projection matrices (W_X, W_Y) , which then are used to obtain the reduced data matrices $Z_X =$

 XW_X , and $Z_Y = YW_Y$.

One of the key differences between PLS and CCA is that while both perform SDR, CCA also simultaneously performs IDR implicitly. Indeed, it involves multiplication of C_{XY} by $C_{XX}^{-1/2}$ on the left and $C_{YY}^{-1/2}$ on the right, which, in turn, requires finding singular values of the X and the Y data matrices independently.

3.1.4 Regularized CCA - rCCA

While CCA is a useful method for finding the maximum correlating features between two sets of data, it does have some limitations. Specifically, in the undersampled regime, where $T \leq \max(N_X, N_Y)$, the matrices C_{XX} and C_{YY} are singular and their inverses do not exist. Using the pseudoinverse to solve the problem can lead to numerical instability and sensitivity to noise. Regularized CCA (rCCA) [47, 52] overcomes this problem by adding a small regularization term to the covariance matrices, allowing them to be invertible. Specifically, one tales

$$\tilde{C}_{XX} = C_{XX} + c_X I_X,\tag{3.6}$$

$$\tilde{C}_{YY} = C_{YY} + c_Y I_Y, \tag{3.7}$$

where \tilde{C}_{XX} , \tilde{C}_{YY} are the new regularized matrices, c_X , $c_Y > 0$ are small regularization parameters and I_X , I_Y are identity matrices with sizes $N_X \times N_X$, $N_Y \times N_Y$ respectively.

This original implementation of rCCA resulted in correlation matrices with diagonals not equal to one. Thus, a better implementation uses a different form of regularization [52] by adding the regularization parameters c_X and c_Y individually to the equations as an affine combination (i. e., $\sum_{i=1}^{n} c_i = 1$) as the following:

$$\tilde{C}_{XX} = \frac{1}{T} (c_{X_1} w_X^{\top} X^{\top} X w_X + c_{X_2} w_X^{\top} w_X)$$
(3.8)

$$\tilde{C}_{YY} = \frac{1}{T} (c_{Y_1} w_Y^{\top} Y^{\top} Y w_Y + c_{Y_2} w_Y^{\top} w_Y). \tag{3.9}$$

This results in the regularized equations for X and Y to be:

$$\tilde{C}_{XX} = \frac{1}{T} \left((1 - c_X) w_X^\top X^\top X w_X + c_X w_X^\top w_X \right) \tag{3.10}$$

$$\tilde{C}_{YY} = \frac{1}{T} ((1 - c_Y) w_Y^{\top} Y^{\top} Y w_Y + c_Y w_Y^{\top} w_Y),$$
 (3.11)

where c_X and c_Y are the regularization parameters, with values between 0 and 1. Writing the regularization conditions in this form is in fact a convex interpolation problem between PLS and CCA, which is a more robust solution and does not suffer from shortening the length of correlations due to the added regularization. As a result, this implementation of rCCA achieves the best accuracy among all other methods.

3.2 Assessing Success and Sampling Noise Treatment

To assess the success of DR, we calculated the ratio between the total correlation between $Z_{X_{\text{test}}}$ and $Z_{Y_{\text{test}}}$, defined as in Eq. (3.2), and the total correlation between X and Y, which we input into the model. Specifically, we take the total correlation as the Frobenius norm of the correlation matrix, $||A||_F = \sqrt{\sum_i \sigma_i^2(A)}$, where $\sigma(A)$ are the singular values of the matrix A. Therefore, the metric of the quality of the DR is

$$\mathcal{RC} = \frac{||\operatorname{Corr}(Z_{X_{\text{test}}}, Z_{Y_{\text{test}}})||_F}{||\operatorname{Corr}(P, P)||_F} = \frac{||\operatorname{Corr}(Z_{X_{\text{test}}}, Z_{Y_{\text{test}}})||_F}{m_{\text{shared}}},$$
(3.12)

where Corr stands for the correlation matrix between its arguments, and we use $||\operatorname{Corr}(P,P)||_F = m_{\text{shared}}$ as the total shared correlation that one needs to recover. Statistical fluctuations aside, \mathcal{RC} should vary between zero (bad reconstruction of the shared variables) and one (perfect reconstruction).

In many real-world applications, the number of available samples, T, is often

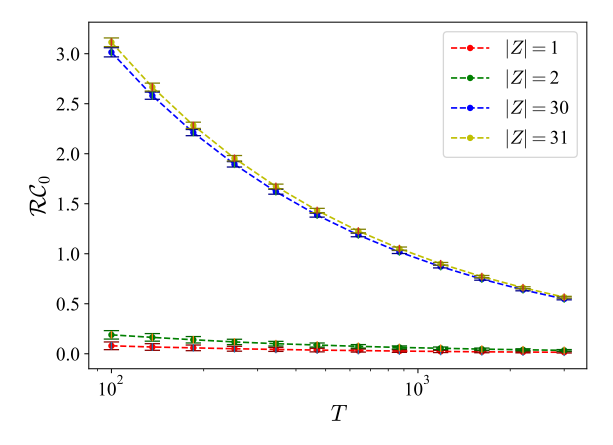


Figure 3.1: The resulting correlations are averages of all the points in the phasespace, then averaged over 10 different realizations of the matrices. The error bars are for two standard deviations around the mean

limited compared to the dimensionality of the data, N_X and N_Y . This undersampling can introduce spurious correlations. We are not aware of analytical results to calculate the effects of the sampling noise on estimating singular values in the model in Eq. (2.1) [5]. Thus, to estimate the effect of the sampling noise, we adopt an empirical approach. Specifically, we generate two random matrices, $Z_{X_{\text{random}}}$ and $Z_{Y_{\text{random}}}$, of sizes $T \times |Z_X|$ and $T \times |Z_Y|$, respectively. We then calculate the correlation between these matrices, denoted as \mathcal{RC}_0 , for multiple such trials using the metric in Eq. (3.12). For random $Z_{X_{\text{random}}}$ and $Z_{Y_{\text{random}}}$, \mathcal{RC} should be zero. However, Fig. 3.1 shows that, especially for large dimensionalities of the compressed variables and small T, the sampling noise results in a significant spurious $\mathcal{RC}_0 > 0$, which may even be larger than 1! Crucially,

 \mathcal{RC}_0 does not fluctuate around its mean across trials, so that the sampling bias is narrowly distributed.

To compensate for this sampling bias, we subtract it from the reconstruction quality metric,

$$\mathcal{RC}' = \mathcal{RC} - \mathcal{RC}_0. \tag{3.13}$$

It is this \mathcal{RC}' that we plot in all Figures in this paper as the ultimate metric of the reconstruction quality. While subtracting the bias is not the most rigorous mathematically, it provides a practical approach for reducing the effects of the sampling noise.

3.3 Implementation

We used Python and the scikit-learn [38] library for performing PCA, PLS, and CCA, while the cca-zoo [7] library was used for rCCA. For PCA, SVD was performed with default parameters. For PLS, the PLS Canonical method was used with the NIPALS algorithm. For both PLS and CCA, the tolerance was set to 10^{-4} with a maximum convergence limit of 5000 iterations. For rCCA, regularization parameters were set as $c_1 = c_2 = 0.1$. All other parameters not explicitly here were set to their default values.

All figures shown in this paper were averaged over 10 independent realizations of R_X , R_Y , U_X , U_Y , P, while fixing the projection matrices V_X , V_Y , Q_X , Q_Y . We then performed an additional round of averaging everything over 10 realizations of the projection matrices themselves. The simulations were parallelized and run on Amazon Web Services (AWS) servers of instance types ml.c5.2xlarge.

Chapter 4

Results

We perform numerical experiments to explore the undersampled regime, $T \lesssim N_X, N_Y$. We use $T = \{100, 300, 1000, 3000\}$ samples, $N_X = N_Y = 1000$. We explore the case of one shared signal only, $m_{\rm shared} = 1$ and we mask this shared signal by a varying number of self signals and noise. We vary the number of retained dimensions, $(|Z_X|, |Z_Y|)$, and explore how many of them are needed to recover the shared signal in the noise and the self signal background with different SNR.

For brevity, we explore two cases: (1) One self-signal in X and Y in addition to the shared signal $(m_{\text{self}} = 1)$; (2) many self-signals in X and Y. For both cases, we calculate the quality of reconstruction as the function of the shared and the self SNR, γ_{shared} and γ_{self} . In all figures, we show \mathcal{RC}' for severely undersampled (first row, T = 300) and relatively well sampled (second row, T = 3000) regimes. We also show the value of \mathcal{RC}_0 , the bias that we removed from our reconstruction quality metric, for completeness, see section 3.2 for details.

4.1 One self-signal in X and Y in addition to the shared signal $(m_{\text{self}} = 1)$

4.1.1 Keeping 1 dimension after reduction $(|Z_{X/Y}| = 1)$

Figure 4.1 shows that, in Case 1, when one dimension is retained in DR of X and Y, PCA populates the compressed variable with the largest variance signals and hence struggles to retain the shared signal when $\gamma_{\text{self}} > \gamma_{\text{shared}}$, regardless of the number of samples. However, both PLS and rCCA excel in achieving nearly perfect reconstructions. When $T \ll N_X$, straightforward CCA cannot be applied (see 3.1.3-3.1.4), but it too achieves a perfect reconstruction when $T > N_X$.

4.1.2 Keeping 2 dimensions after reduction $(|Z_{X/Y}| = 2)$

In Fig. 4.2, we allow two dimensions in the reduced variables. For PCA, we expect this to be sufficient to preserve both the self and the shared signals. Indeed, PCA now works for all γ s and T, although with a slightly reduced accuracy for large shared signals compared to Fig. 4.1. PLS and rCCA continue to deliver highly accurate reconstructions. So does the CCA for $T > N_X$. Spurious correlations, as measured by \mathcal{RC}_0 grow slightly with the increasing dimensionality of Z_X , Z_Y compared to Fig. 4.1. This is expected since more projections must now be inferred from the same amount of data.

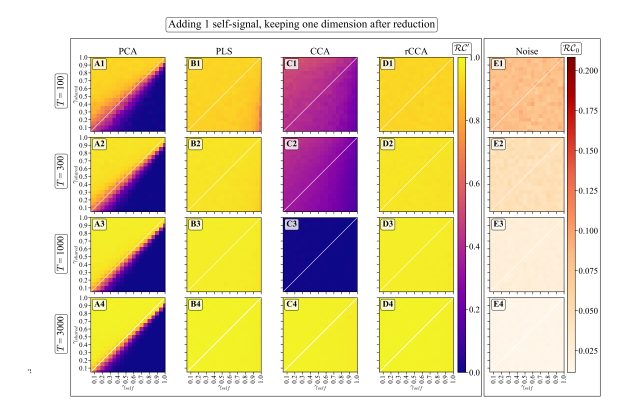


Figure 4.1: Performance of PCA, PLS, CCA, rCCA, and noise in recovery of the shared signal for $|Z_X| = |Z_Y| = 1 = m_{\rm self}$. PCA struggles to detect shared signals when they are weaker than the self signals. PLS and rCCA demonstrate nearly perfect reconstruction. CCA displays no reconstruction in the undersampled regime $T \ll N_X$, and it is nearly perfect for large T.

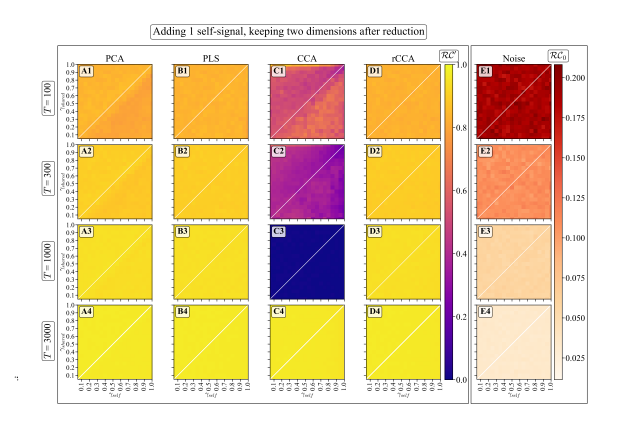


Figure 4.2: Same as Fig. 4.1, but for $|Z_X| = |Z_Y| = 2 = m_{\text{self}} + m_{\text{shared}}$. Now there are enough compressed variables for PCA to detect the shared signal. Other methods perform similarly to Fig. 4.1, albeit the noise is larger.

4.2 Many self-signal in X and Y in addition to the shared signal $(m_{self} = 30)$

4.2.1 Keeping 1 dimension after reduction $(|Z_{X/Y}| = 1)$

We now turn to $m_{\rm self} \gg m_{\rm shared}$. We use $m_{\rm shared} = 1$, $m_{\rm self} = 30$ for concreteness. We expect that the performance of SDR methods will degrade weakly, as they are designed to be less sensitive to the masking effects of the self signals. In contrast, we expect IDR to be more easily confused by the many strong self-signals, degrading the performance. Indeed, Fig. 4.3 shows that PCA now faces challenges in detecting shared signals, even when the self signals are weaker than in Fig. 4.1. Increasing T improves its performance only slightly. Somewhat surprisingly, PLS performance also degrades, with improvements at $T \gg N_X$. CCA again displays no reconstruction when $T \ll N_X$, switching to near perfect reconstruction at large T. Crucially, rCCA again shines, maintaining its strong performance, consistently demonstrating nearly perfect reconstruction.

4.2.2 Keeping 30 dimensions after reduction ($|Z_{X/Y}| = 30$)

Since one retained dimension is not sufficient for PCA to represent the shared signal when $\gamma_{\text{shared}} \lesssim \gamma_{\text{self}}$, we increase the dimensionality of reduced variables $|Z_X| = |Z_Y| = m_{\text{self}} \gg m_{\text{shared}}$), cf. Fig. 4.4. PCA now detects shared signals even when they are weaker than the self-signals, $\gamma_{\text{shared}} < \gamma_{\text{self}}$, but at a cost of the reconstruction accuracy plateauing significantly below 1. In other words, when self and shared signals are comparable, they mix, allowing for partial reconstruction. However, even at $T \gg N_X$, PCA cannot break into the phase diagram's lower right corner. Other methods perform similarly, reconstructing shared signals over the same or wider ranges of sampling and the SNR ratios than in Fig. 4.3. For all of them, the improvement comes

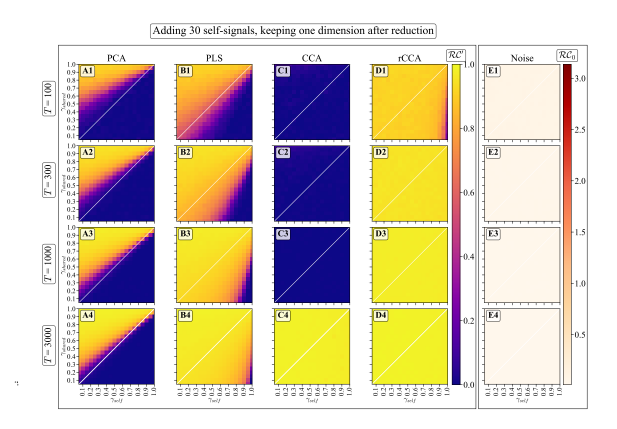


Figure 4.3: Reconstruction results for $m_{\text{self}} = 30$, $m_{\text{shared}} = 1$, and $|Z_X| = |Z_Y| = 1$. PCA struggles to detect any shared signals when they are even comparable to the self ones. PLS performance also degrades. CCA displays its usual impotence at small T. Finally, rCCA demonstrates nearly perfect reconstruction for all parameter values.

at the cost of the decreased asymptotic performance. The most distinct feature of this regime is the dramatic effect of noise, where 30-dimensional compressed variables can accumulate enough sampling fluctuations to recover correlations that are supposedly nearly twice as high as the data actually has.

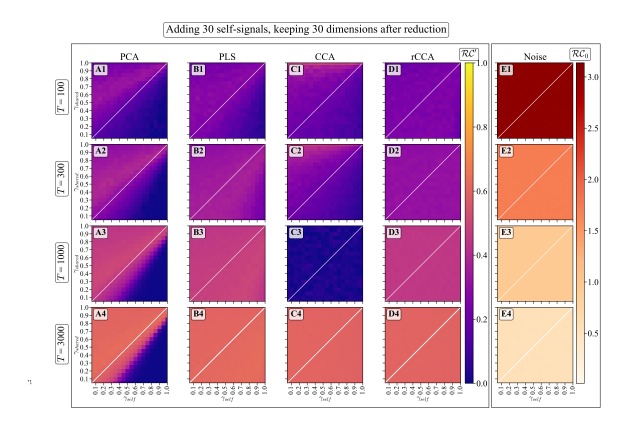


Figure 4.4: DR performance for $|Z_X| = |Z_Y| = m_{\rm self} > m_{\rm shared}$). PCA now detects shared signals even when they are weaker than the self signals. However, the quality of reconstruction is significantly lower than in Fig. 4.2. PLS detects signals in a larger part of the phase space, but also with a significant reduction in quality, which improves with sampling. CCA has its usual problem for $T \ll N_X$, and, like PLS, it has a significantly lower reconstruction quality than in the regime in Fig. 4.3. rCCA is able to detect the signal in the whole phase space, but again with worse quality. Finally, spurious correlations are high, though they decrease with better sampling.

4.2.3 Keeping 31 dimensions after reduction ($|Z_{X/Y}| = 31$)

Figure 4.5 now explores a regime when the dimensionality of the compressed variables is enough to store both the self and the shared interactions at the same time, $|Z_X| = |Z_Y| = m_{\text{self}} + m_{\text{shared}} = 31$. With just one more dimension than Fig. 4.4, PCA abruptly transitions to being able to recover shared signals for all SNRs, albeit still saturating at a far from perfect performance at large T. PLS, CCA, rCCA, and noise show behavior remain similar to Fig. 4.4.

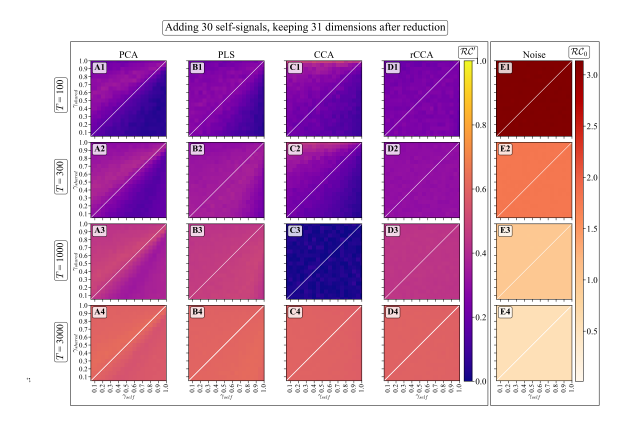


Figure 4.5: PCA, PLS, CCA, rCCA, and noise results when 31 dimensions are kept after reduction ($|Z_X| = |Z_Y| = m_{\text{self}} + m_{\text{shared}}$). PCA now can detect more shared signals when they are weaker than the self signals (A1), however, with a significantly lower quality compared to figure 4.2, but suddenly explores the whole phase space, still with lower accuracy than Case 1. PLS, CCA, rCCA, and noise show similar behavior to figure 4.4.

4.3 Key Parameters and Testing Technique for Dimensionality of Self and Shared Signals

Our analysis suggests that there are three relevant factors that determine the ability of DR to reconstruct shared signals. The first is the strength of the shared and the self signals compared to each other and to noise. For brevity, in the following analysis, we fix γ_{self} and define the ratio $\tilde{\gamma} = \gamma_{\text{shared}}/\gamma_{\text{self}}$ to represent this effect. The second factor affecting the performance is the ratio between the number of shared and self signals, denoted by $\tilde{m} = m_{\text{shared}}/m_{\text{self}}$. The third factor is the number of samples per dimension of the reduced variable, denoted by $\tilde{q} = T/|Z|$.

In Fig. 4.6, we illustrate how these parameters influence the performance of DR, \mathcal{RC}' . Each subplot varies \tilde{q} , while holding T constant and changing $|Z_X|$. We compare the results of PCA (representing IDR) and rCCA (representing SDR). Each curve is averaged over 10 trials, with error bars indicating 1 standard deviation around the mean, using algorithmic parameters as described in section 3.3.

We see that the relative strength of signals, as represented by $\tilde{\gamma}$, plays a significant role in determining which method performs better. If the shared signals are larger (bottom) both approaches work. However, for weak shared signals (top), SDR is generally more effective. Further, the ratio between the number of shared and self signals, \tilde{m} , also plays an important role. When \tilde{m} is large (left), IDR is more likely to detect the shared signal before the self signals, and it approaches the performance of SDR. However, when \tilde{m} is small, IDR is more likely to capture the self signals before moving on to the shared signals, degrading performance (right). Finally, not surprisingly, the number of samples per dimension of the compressed variables, \tilde{q} , is also critical to the success. If \tilde{q} is small, the signal is drowned in the sampling noise, and adding more retained dimensions hurts the DR process. This expresses itself as a peak for SDR performance around $|Z_X| = m_{\rm shared}$. For IDR, the peak is around

 $|Z_X| = m_{\text{self}} + m_{\text{shared}}$, thus requiring more data to achieve performance similar to SDR.

We observe that the performance of rCCA (SDR) is almost independent of changing \tilde{m} or $\tilde{\gamma}$, indicating that it focuses on shared dimensions even if the latter is masked by self signals. The algorithm crucially depends on \tilde{q} , where adding more dimensions (decreasing \tilde{q}) than needed hurts the reduction. This is because, for a fixed number of samples, the reconstruction of each dimension then gets worse. In contrast, for PCA (IDR), the performance depends on all three relevant parameters, \tilde{q} , \tilde{m} , and $\tilde{\gamma}$. At some parameter combinations, the performance of IDR in reconstructing shared signals approaches SDR. However, in all cases, SDR never performs worse than IDR on this task.

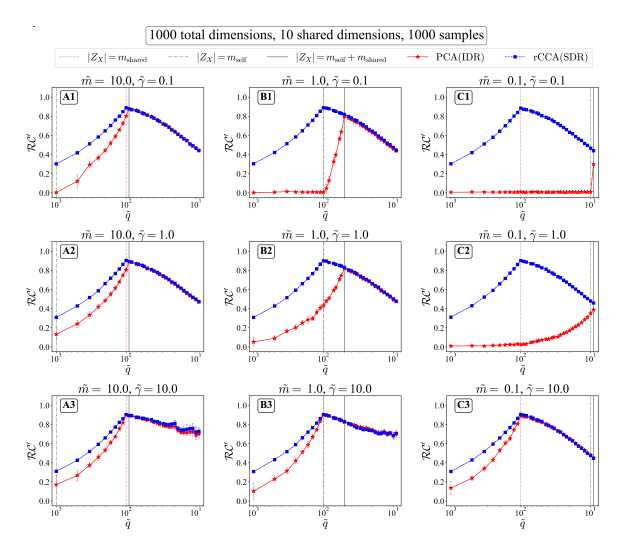


Figure 4.6: Performance of PCA (IDR) and rCCA (SDR) for different values of the relevant parameters of the model: the number of samples per dimension of the compressed variable (\tilde{q}) , the strength of shared signals relative to the self ones $(\tilde{\gamma})$, and the ratio of the number of shared to self signal components (\tilde{m}) , while fixing the number of samples (T=1000) and the number of shared dimensions $(m_{\text{shared}}=10)$. Note that decreasing \tilde{q} (left to right) corresponds to increasing the dimension of the latent space $|Z_X|$ at a fixed number of samples T.

Chapter 5

Discussions

5.1 Extensions and Generalizations

We proposed a generative linear model, which we believe captures multiple desired features of multimodal data with shared and non-shared signals. The model focused only on data with two measured modalities. However, while not a part of this study, the model can be readily extended to accommodate more than two modalities (e. g., $X_i = R_i + U_i V_i + PQ_i$ for i = 1, ..., n, where n represents the number of modalities). Then, methods such as Tensor CCA, which can handle more than two modalities [32], can be used to get insight into DR on such data.

5.2 Explaining Observations in the Literature

We analyzed different DR methods on data generated by this model in different parameter regimes. We showed that linear SDR methods are clearly superior to their IDR counterparts for detecting shared signals. We can thus make a strong practical suggestion that, whenever the goal is to reconstruct a low dimensional representation of covariation between two components of the data, IDR methods (PCA) should always be avoided in favor of SDR. Of the examined SDR approaches, rCCA is a

clear winner in all parameter regimes and should always be preferred. This finding explains the results of, for example, [41] and others that SDR can recover joint neuro-behavioral latent spaces with fewer latent dimensions and using fewer samples than IDR methods. Further, our observation that SDR is always superior to IDR in the context of our model corroborates the theoretical findings of [34], who proved a similar result in the context of discrete data and a different SDR algorithm, namely the Symmetric Information Bottleneck [16]. Collectively, these diverse investigations, linear and nonlinear, theoretical, computational, and empirical, provide strong evidence that generic (not just linear) SDR methods are likely to be more efficient in extracting covariation than their IDR analogs.

5.3 Is SDR strictly effective in low sampling situations?

Our study answers an open question in the literature surrounding the effectiveness of SDR techniques. Specifically, there has been debate about whether PLS, an SDR method, is effective at low sampling [9, 21, 17, 18]. Our results show that SDR is not necessarily effective in the undersampled regime. It works well when the number of samples per retained dimension is high (even if the number of samples per observed dimension is low), but only when the dimensionality of the reduced description is matched to the actual dimensionality of the shared signals.

5.4 Diagnostic Test for number of latent signals

In addition to the previous, our results can be used as a diagnostic test to determine the number of shared versus self signals in data. As demonstrated in Fig. 4.6, total correlations between Z_X and Z_Y obtained by applying PCA and rCCA increase monotonically as the dimensionality of Zs increases, until this dimensionality becomes larger than the signal dimensionality. For PCA, the signal dimensionality is equal to the sum of the number of the shared and the self signals, $m_{\rm shared} + m_{\rm self}$. For rCCA, it is only the number of the shared signal. Thus increasing the dimensionality of the compressed variables and tracking the performance of rCCA and PCA until they diverge can be used to identify the number of self signals in the data, provided that the data, indeed, has a low-dimensional latent structure. This approach can be a valuable tool in various applications, where the characterization of shared and self signals in complex systems can provide insights into their structure and function.

5.5 Limitations, and Future Work

5.5.1 Linearity of the model

While this work has provided useful insight, the assumptions made here may not fully capture the complexity of real-world data. Specifically, our data is generated by a linear model with random Gaussian features. It is unlikely that real data have this exact structure. Therefore, there is a need for further exploration of the advantages and limitations of linear DR methods on data that have a low-dimensional, but nonlinear shared structure. This can be done using more complex nonlinear generative models, such as nonlinearly transforming the data generated by Eq. (2.1-2.2), or random feature two-layered neural network models [39].

5.5.2 Linearity of the methods

A different possible future research direction is to explore the performance of non-linear DR methods on data from generative models with a latent low-dimensional nonlinear structure. Autoencoders and their variational extensions are a natural extension of IDR to learn nonlinear reduced dimensional representations [24, 28, 23].

Meanwhile, Deep CCA and its variational extensions [1, 49, 6, 48] should be explored as a nonlinear version of SDR. Both of these types of methods can potentially capture more complex relationships between the modalities and improve the quality of the reduced representations, and it is not clear if the SDR class of methods is always more efficient than the IDR one.

5.5.3 Linearity of the metric

Our analysis also depends on the choice of metric used to quantify the performance of DR, and different choices should also be explored. For example, to capture nonlinear correlations, mutual information can be utilized to quantify the relationships between the reduced representations.

5.6 Conclusion

In conclusion, we highlight a general principle that, when searching for a shared signal between different modalities of data, SDR methods are preferable to IDR methods. Additionally, the differences in performance between the two classes of methods can tell us a lot about the underlying structure of the data. Finally, for a limited number of samples, naive approaches, such as increasing the number of compressed dimensions indefinitely to overcome the masking of shared signals by self signals are infeasible. Thus, the use of SDR methods becomes even more essential in such cases, and despite the aforementioned limitations, we believe that our work provides a compelling addition to the body of knowledge that SDR outperforms IDR in detecting shared signals quite generally.

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