

Principal Component Analysis

Dimensionality Reduction and the Maximum Variance Perspective

Based on Bishop PRML Chapter 12

November 9, 2025

The Dimensionality Challenge

Modern machine learning confronts us with high-dimensional data:

- ▶ Images: 28×28 grayscale = 784 dimensions
- ▶ Genomics: thousands of gene expressions per sample
- ▶ Text: vocabulary size in tens of thousands
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Problems we face:

- ▶ Cannot visualize beyond 3D
- ▶ Computational cost scales poorly with dimension
- ▶ Many algorithms struggle in high dimensions (curse of dimensionality)

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The central question: What does “meaningful” mean mathematically?

Real-World Applications

figures/eigenfaces.png

(a) Face recognition using eigenfaces

figures/genomics_pca.png

(b) Gene expression analysis

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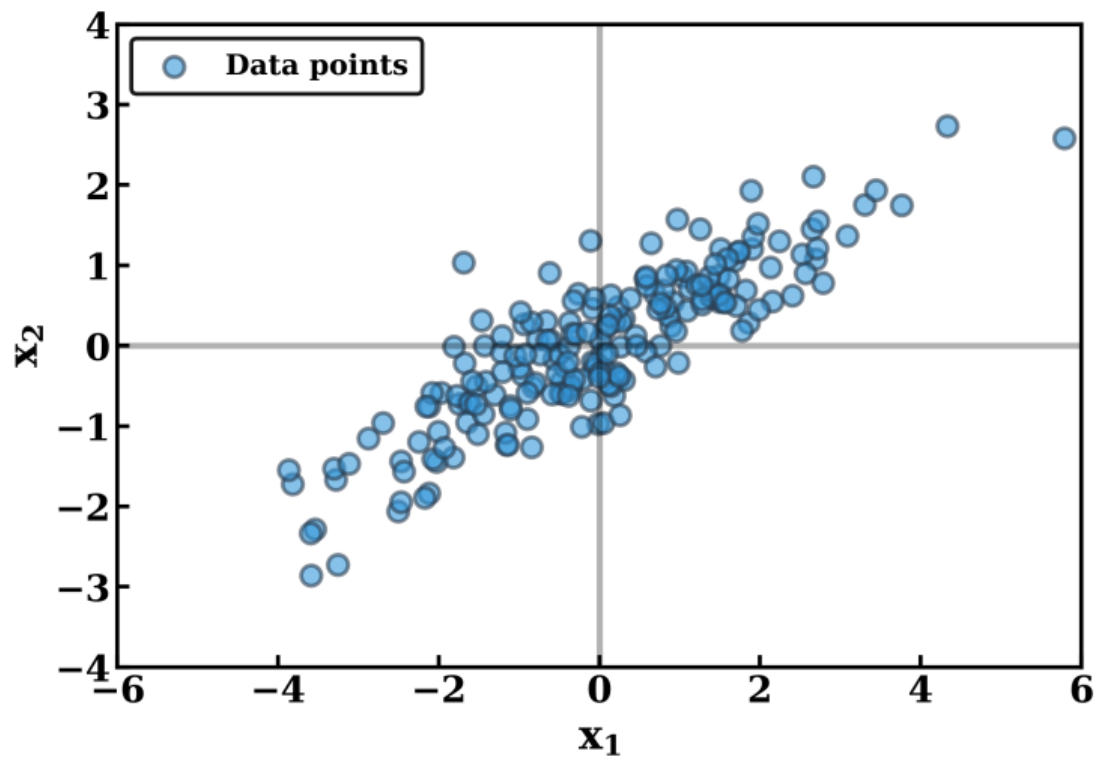
Which projection is “best”?

Principal Component Analysis provides an answer based on two equivalent perspectives:

1. **Maximum variance:** Keep directions with highest variance
2. **Minimum error:** Minimize information loss from projection

Both lead to the same solution.

A Simple 2D Example



2D data points forming an elliptical cloud

Consider data naturally lying along an elongated cloud. There's clearly more "spread"

Intuition: Direction of Spread

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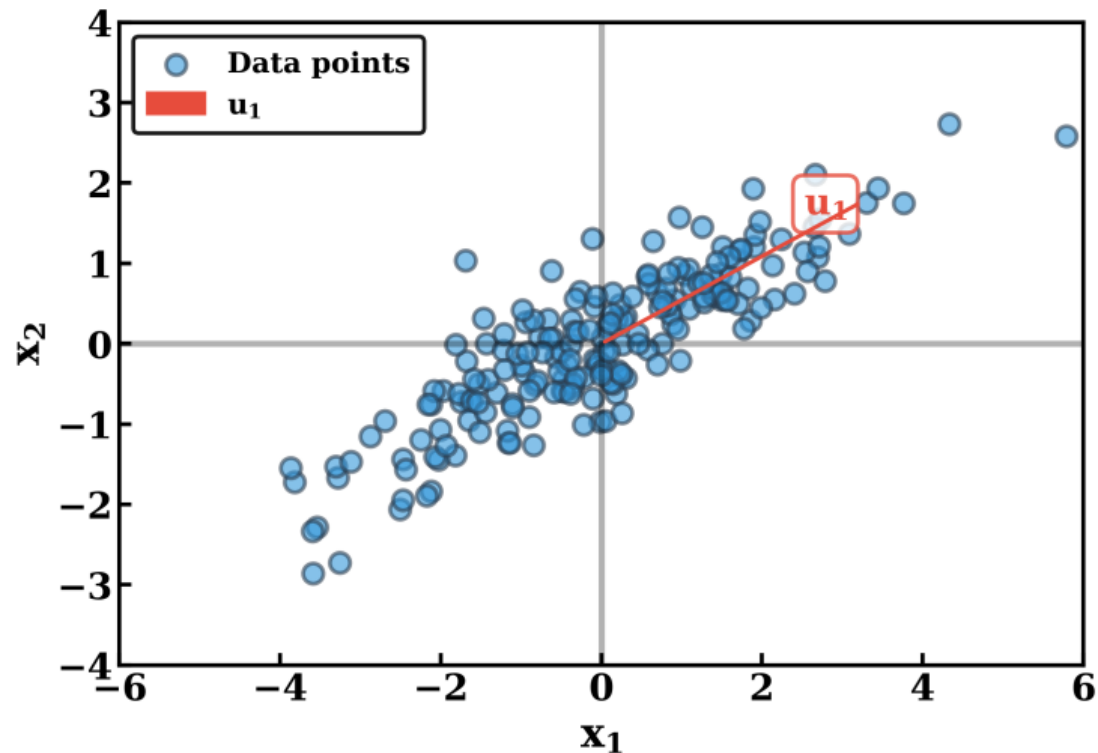
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Key insight: Directions with high variance contain more “information” about the structure of the data.

If we must reduce from 2D to 1D, we should keep the direction along which the data varies most.

This direction of maximum variance is the **first principal component**.

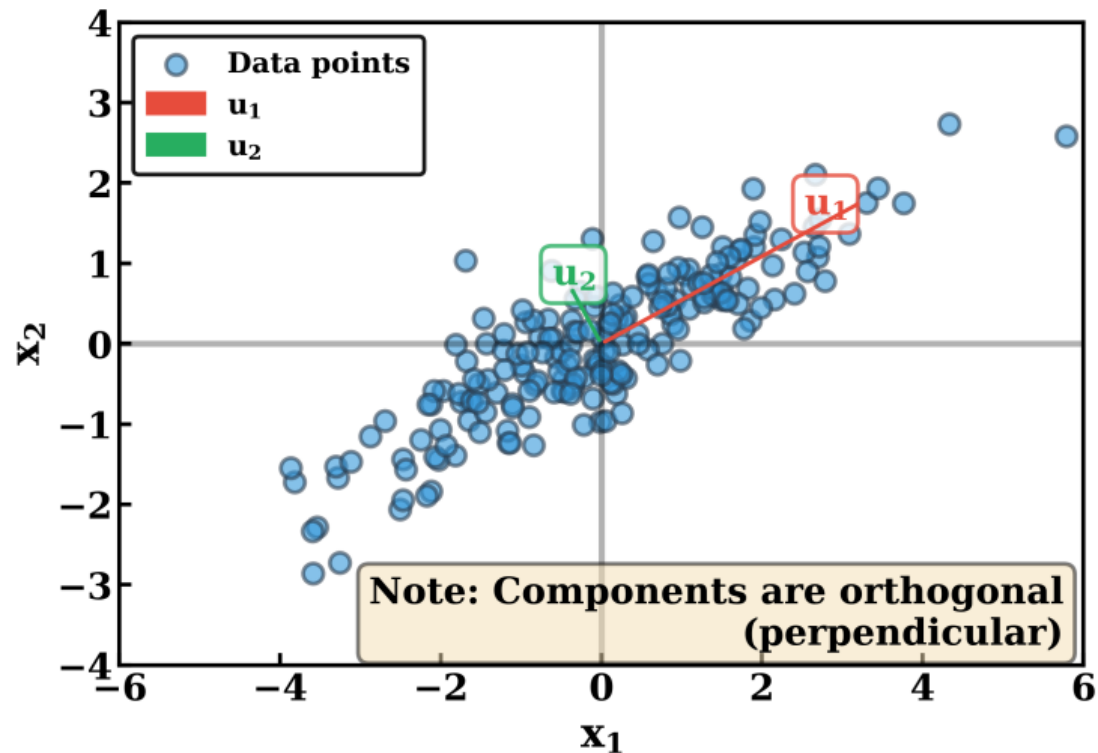
Visual: First Principal Component



First principal component \mathbf{u}_1 captures maximum variance

The first principal component \mathbf{u}_1 points in the direction of maximum variance.

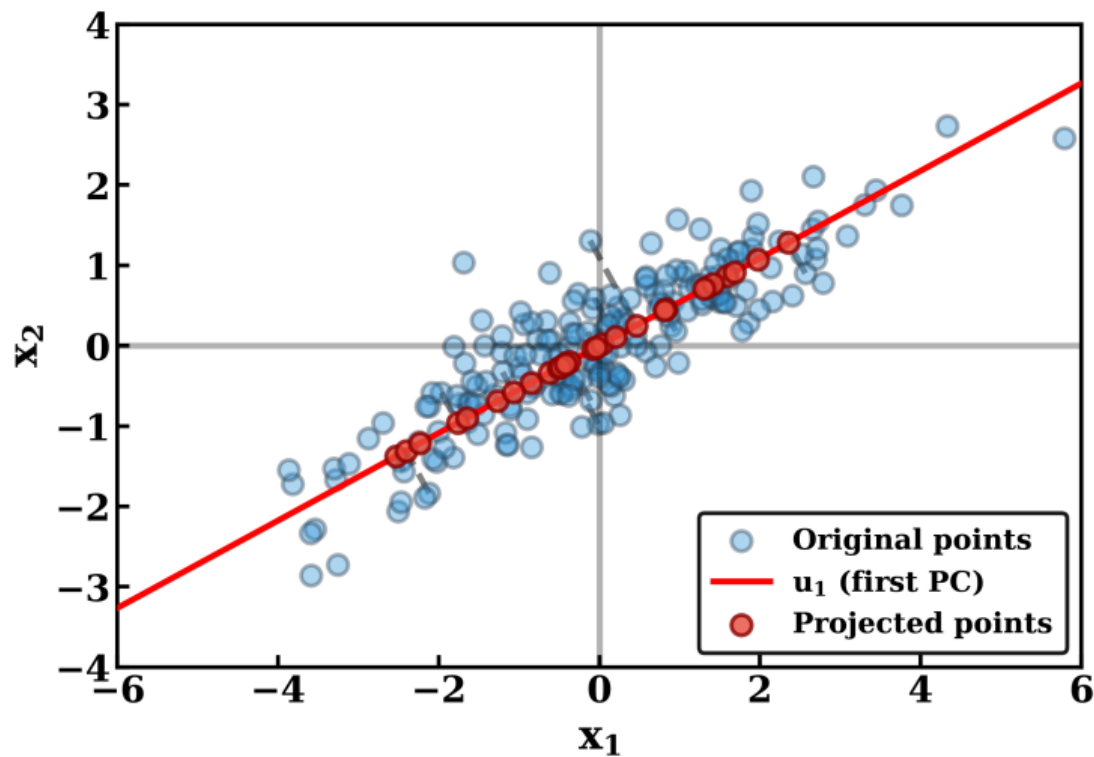
Visual: Second Principal Component



Both principal components shown as orthogonal vectors

The second principal component u_2 is orthogonal to the first and captures the

Projection Intuition



Projection onto first principal component reduces dimensionality

Projecting onto u_1 reduces dimensionality from 2D to 1D while preserving maximum variance.

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Reduced data: 1 dimension (coordinate along \mathbf{u}_1)

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What did we keep?

- ▶ Variance in the principal direction
- ▶ This is the “large” variance—more important
- ▶ The overall structure and spread of the data

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For dimensionality reduction: project data onto the top M principal components.

This captures as much variance as possible in M dimensions.

Setup: The Data Matrix

Consider a dataset of N observations, each in D dimensions:

$$\mathbf{x}_1, \mathbf{x}_2, \dots, \mathbf{x}_N \quad \text{where } \mathbf{x}_n \in \mathbb{R}^D$$

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Why centering matters: PCA finds directions of maximum variance around the mean. Without centering, we'd be finding variance around the origin, which is not meaningful.

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The projection of each data point $\tilde{\mathbf{x}}_n$ onto \mathbf{u}_1 is:

$$\mathbf{u}_1^T \tilde{\mathbf{x}}_n$$

This is a scalar—the coordinate along \mathbf{u}_1 .

Variance of Projections

The projected data are the scalars: $\mathbf{u}_1^T \tilde{\mathbf{x}}_1, \mathbf{u}_1^T \tilde{\mathbf{x}}_2, \dots, \mathbf{u}_1^T \tilde{\mathbf{x}}_N$

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Since the data are mean-centered, the mean of projections is zero:

$$\frac{1}{N} \sum_{n=1}^N \mathbf{u}_1^T \tilde{\mathbf{x}}_n = \mathbf{u}_1^T \left(\frac{1}{N} \sum_{n=1}^N \tilde{\mathbf{x}}_n \right) = \mathbf{u}_1^T \mathbf{0} = 0$$

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The sample variance of projections is:

$$\frac{1}{N} \sum_{n=1}^N (\mathbf{u}_1^T \tilde{\mathbf{x}}_n)^2$$

Goal: Maximize this quantity over choice of \mathbf{u}_1 .

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Note: \mathbf{S} is symmetric and positive semi-definite.

Variance in Matrix Form

The variance of projections can now be written as:

$$\begin{aligned}\frac{1}{N} \sum_{n=1}^N (\mathbf{u}_1^T \tilde{\mathbf{x}}_n)^2 &= \frac{1}{N} \sum_{n=1}^N \mathbf{u}_1^T \tilde{\mathbf{x}}_n \tilde{\mathbf{x}}_n^T \mathbf{u}_1 \\ &= \mathbf{u}_1^T \left(\frac{1}{N} \sum_{n=1}^N \tilde{\mathbf{x}}_n \tilde{\mathbf{x}}_n^T \right) \mathbf{u}_1 \\ &= \mathbf{u}_1^T \mathbf{S} \mathbf{u}_1\end{aligned}$$

Our optimization problem:

$$\max_{\mathbf{u}_1} \mathbf{u}_1^T \mathbf{S} \mathbf{u}_1 \quad \text{subject to} \quad \mathbf{u}_1^T \mathbf{u}_1 = 1$$

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Form the Lagrangian:

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Taking the derivative with respect to \mathbf{u}_1 and setting to zero:

$$\frac{\partial \mathcal{L}}{\partial \mathbf{u}_1} = 2\mathbf{S}\mathbf{u}_1 - 2\lambda_1\mathbf{u}_1 = \mathbf{0}$$

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This is an eigenvalue equation!

- ▶ \mathbf{u}_1 is an eigenvector of the covariance matrix \mathbf{S}
- ▶ λ_1 is the corresponding eigenvalue

See Bishop eq. 12.6

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Conclusion: The variance captured equals the eigenvalue λ_1 .

To maximize variance, choose the eigenvector with the **largest eigenvalue**.

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where \mathbf{u}_2 is the eigenvector with the second-largest eigenvalue λ_2 .

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General pattern: The i -th principal component is the eigenvector corresponding to the i -th largest eigenvalue.

Complete Solution

The covariance matrix has eigendecomposition:

$$\mathbf{S} = \sum_{i=1}^D \lambda_i \mathbf{u}_i \mathbf{u}_i^T$$

where $\lambda_1 \geq \lambda_2 \geq \dots \geq \lambda_D \geq 0$.

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- ▶ $\mathbf{u}_1, \mathbf{u}_2, \dots, \mathbf{u}_D$ (eigenvectors)
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The variance captured by the i -th component is λ_i .

Summary of Maximum Variance Formulation

Problem: Find low-dimensional projection preserving maximum variance.

Solution:

1. Compute the sample covariance matrix \mathbf{S}
2. Find eigenvalues and eigenvectors of \mathbf{S}
3. Sort eigenvalues: $\lambda_1 \geq \lambda_2 \geq \dots \geq \lambda_D$
4. The first M principal components are the eigenvectors $\mathbf{u}_1, \dots, \mathbf{u}_M$
5. Project data: $\mathbf{z}_n = \mathbf{U}_M^T \tilde{\mathbf{x}}_n$ where $\mathbf{U}_M = [\mathbf{u}_1 \cdots \mathbf{u}_M]$

Total variance captured: $\sum_{i=1}^M \lambda_i$

Total variance in data: $\sum_{i=1}^D \lambda_i = \text{tr}(\mathbf{S})$

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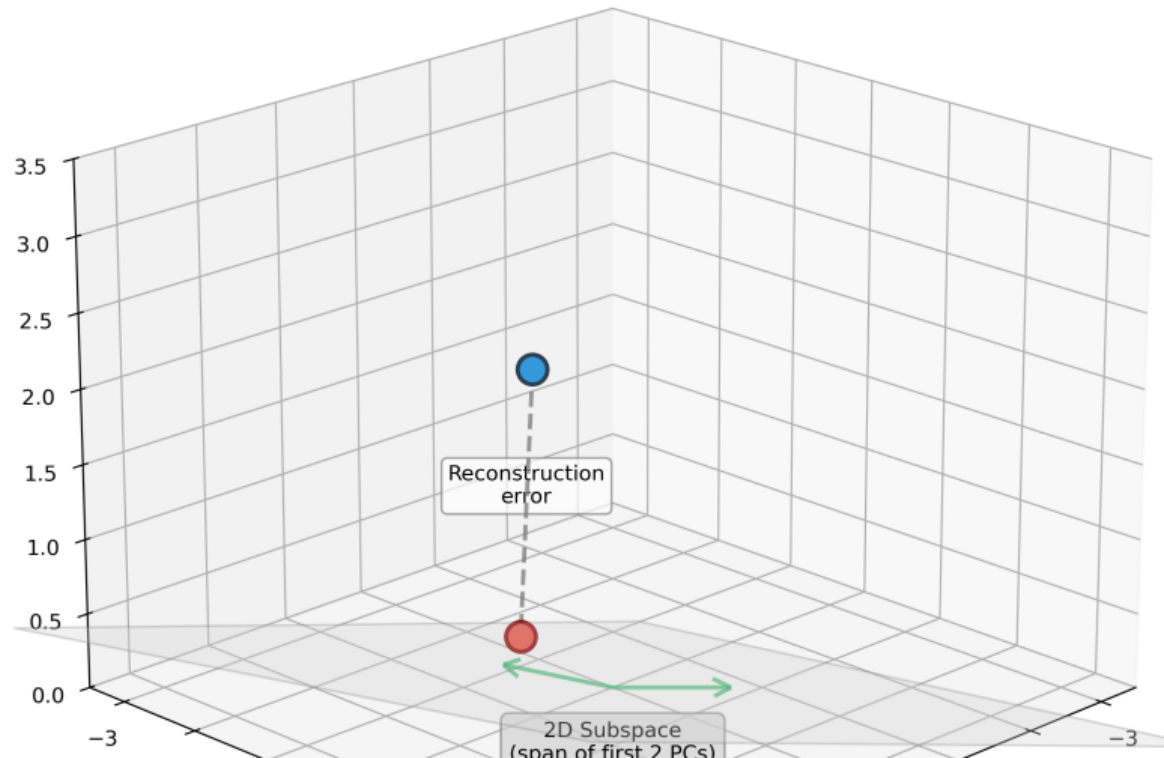
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Idea:

- ▶ Project data onto M -dimensional subspace
- ▶ Reconstruct back to original D dimensions
- ▶ Measure error between original and reconstruction
- ▶ Choose subspace that minimizes this error

The Projection Operation

- Original $\tilde{\mathbf{x}}_n$
- Reconstructed $\hat{\mathbf{x}}_n$



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Reconstruction back to D dimensions:

$$\hat{\mathbf{x}}_n = \mathbf{U}_M \mathbf{z}_n = \mathbf{U}_M \mathbf{U}_M^T \tilde{\mathbf{x}}_n$$

(Note: $\mathbf{U}_M \mathbf{U}_M^T$ is the projection matrix onto the subspace.)

Reconstruction Error

The reconstruction error for point n is:

$$\|\tilde{\mathbf{x}}_n - \hat{\mathbf{x}}_n\|^2 = \|\tilde{\mathbf{x}}_n - \mathbf{U}_M \mathbf{U}_M^T \tilde{\mathbf{x}}_n\|^2$$

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The total reconstruction error across all data points:

$$J = \frac{1}{N} \sum_{n=1}^N \|\tilde{\mathbf{x}}_n - \mathbf{U}_M \mathbf{U}_M^T \tilde{\mathbf{x}}_n\|^2$$

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Goal: Choose the M -dimensional subspace (i.e., choose \mathbf{U}_M) to minimize J .

Equivalence to Maximum Variance

It can be shown that:

$$\sum_{n=1}^N \|\tilde{\mathbf{x}}_n - \hat{\mathbf{x}}_n\|^2 + \sum_{n=1}^N \|\hat{\mathbf{x}}_n\|^2 = \sum_{n=1}^N \|\tilde{\mathbf{x}}_n\|^2 = \text{constant}$$

Equivalence to Maximum Variance

It can be shown that:

$$\sum_{n=1}^N \|\tilde{\mathbf{x}}_n - \hat{\mathbf{x}}_n\|^2 + \sum_{n=1}^N \|\hat{\mathbf{x}}_n\|^2 = \sum_{n=1}^N \|\tilde{\mathbf{x}}_n\|^2 = \text{constant}$$

The first term is variance *lost* (reconstruction error).

The second term is variance *captured* (in the subspace).

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Therefore:

Minimizing reconstruction error \iff Maximizing captured variance

Both formulations yield the same solution! See Bishop §12.1.1

Intuitive Connection

Variance decomposition

Total variance = Captured variance + Lost variance

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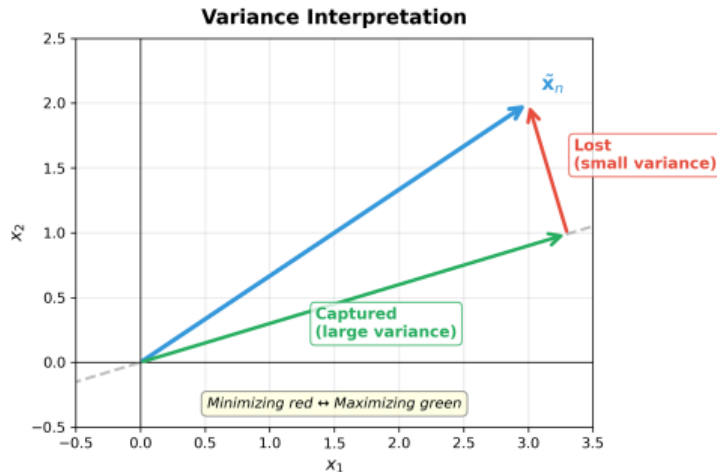
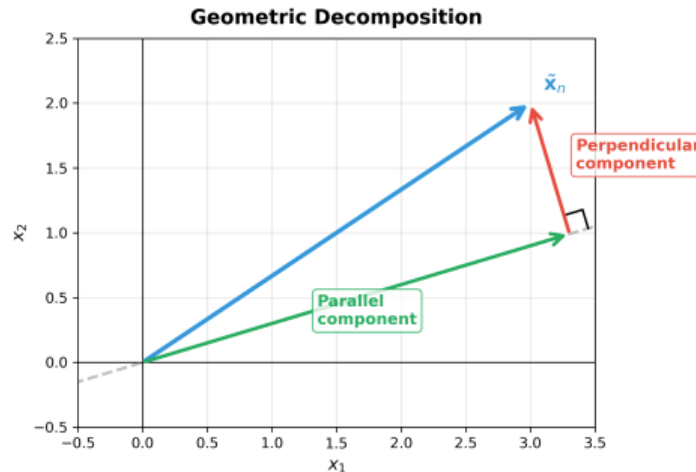
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- ▶ Maximum variance formulation \Leftrightarrow Minimum error formulation

Both perspectives lead to the same principal components.

Geometric Interpretation



Orthogonal decomposition showing variance-error equivalence

The PCA Algorithm

Input: Data matrix \mathbf{X} of size $N \times D$, desired dimension M

Algorithm:

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7. Project: $\mathbf{z}_n = \mathbf{U}_M^T \tilde{\mathbf{x}}_n$ for all n

Output: Low-dimensional representations $\mathbf{z}_1, \dots, \mathbf{z}_N$

Computational Considerations

Covariance matrix: $D \times D$ (can be very large!)

- ▶ For $D = 10,000$ features, \mathbf{S} requires ~ 800 MB (double precision)
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- ▶ Apply SVD directly to centered data matrix $\tilde{\mathbf{X}}$ (size $N \times D$)
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Standard implementations (scikit-learn, R, MATLAB) use SVD internally.

How Many Components?

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No single correct answer—depends on application:

- ▶ Visualization: $M = 2$ or $M = 3$
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Several diagnostic tools help us decide:

1. Scree plot (eigenvalue spectrum)
2. Cumulative variance explained
3. Cross-validation on downstream task
4. Domain knowledge

Scree Plot

[IMAGE PLACEHOLDER]

Description: A scree plot showing eigenvalue magnitude vs. component number. X-axis: “Principal Component” numbered 1 to 20. Y-axis: “Eigenvalue λ_i ;” (log scale preferred). The plot should show a characteristic “elbow” pattern: first few eigenvalues are large and drop rapidly, then there’s an elbow around component 4-5, after which eigenvalues decrease slowly. Mark the elbow point with a dashed vertical line and annotation “Elbow suggests $M \approx 4$ ”. Use blue circles connected by lines. Include a horizontal red dashed line showing “noise floor” for the smallest eigenvalues.

Look for the “elbow” where eigenvalues drop off sharply, then plateau.

Cumulative Variance Explained

[IMAGE PLACEHOLDER]

Description: Plot of cumulative proportion of variance explained vs. number of components. X-axis: “Number of Components M ” from 1 to 20. Y-axis: “Cumulative Variance Explained” from 0 to 1 (or 0% to 100%). Show a monotonically increasing curve that starts steep and gradually flattens. Add horizontal dashed lines at 0.90, 0.95, and 0.99 with labels. Add vertical dashed lines showing how many components needed to reach each threshold (e.g., $M = 5$ for 90%, $M = 8$ for 95%, $M = 12$ for 99%). Use a smooth blue curve.

Proportion of variance explained by first M components:

$$\frac{\sum_{i=1}^M \lambda_i}{\sum_{i=1}^D \lambda_i}$$

Common thresholds: 90%, 95%, or 99% variance explained.

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- ▶ Feature with large variance (e.g., income in dollars) will dominate
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Standard practice: Standardize features to unit variance

$$x_{nd} \leftarrow \frac{x_{nd} - \bar{x}_d}{\sigma_d}$$

where σ_d is the standard deviation of feature d .

This ensures all features contribute on equal footing.

Practical Example Setup

We'll work through PCA on a classic dataset.

Iris Dataset:

- ▶ 150 samples of iris flowers
- ▶ 4 features: sepal length, sepal width, petal length, petal width
- ▶ 3 species: setosa, versicolor, virginica

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This is a perfect example because:

- ▶ Low enough dimension to understand fully
- ▶ Real biological data with known structure
- ▶ Widely recognized in ML community

Example: Original High-Dimensional Data

[IMAGE PLACEHOLDER]

Description: A 2×2 grid of scatter plots showing pairwise relationships between the 4 iris features. Each subplot shows two features against each other (sepal length vs sepal width, sepal length vs petal length, etc.). Points should be colored by species: red for setosa, green for versicolor, blue for virginica. This creates a “pairs plot” or “scatter plot matrix.” Label axes clearly with feature names. Show that in the original feature space, species have some separation but overlap exists in certain views.

The original 4D space shown as pairwise projections. Some separation between species is visible, but overlapping.

Covariance Structure

[IMAGE PLACEHOLDER]

Description: Heatmap of the 4×4 covariance matrix. Rows and columns labeled: sepal length, sepal width, petal length, petal width. Use a diverging colormap (e.g., blue for negative, white for zero, red for positive correlations). Strong positive correlations should appear between petal length/width and sepal length. Sepal width might show weaker or slightly negative correlation with others. Add a colorbar. Annotate cells with numerical correlation values.

The covariance matrix reveals strong correlations between certain features (e.g., petal dimensions).

Eigenvalue Spectrum

[IMAGE PLACEHOLDER]

Description: Scree plot for the Iris dataset showing 4 eigenvalues. X-axis: Component number (1-4). Y-axis: Eigenvalue magnitude. Show bars or connected points. First eigenvalue should be dominant (much larger than others), second eigenvalue moderate, third and fourth small. Include numerical labels on each point showing the actual eigenvalue. Below or beside, show the variance explained by each component as percentages (e.g., PC1: 72.9%, PC2: 22.8%, PC3: 3.7%, PC4: 0.5% — these are approximate typical values for scaled Iris data).

First two components capture ~96% of variance—excellent for 2D visualization!

First Two Principal Components

[IMAGE PLACEHOLDER]

Description: 2D scatter plot of the Iris data projected onto the first two principal components. X-axis: "First Principal Component (PC1)" ranging approximately -3 to 3. Y-axis: "Second Principal Component (PC2)" ranging approximately -2 to 2. Color points by species (same colors as before: red setosa, green versicolor, blue virginica). Show that setosa is clearly separated from the other two species, while versicolor and virginica have slight overlap. This demonstrates that PCA preserves the species structure well. Add a legend showing species colors.

The 2D projection preserves species separation well. Setosa is distinctly separated; some overlap between versicolor and virginica.

Reconstruction Quality

[IMAGE PLACEHOLDER]

Description: Show reconstruction quality comparison. Create a 2×3 grid:

- ▶ Row 1: Three iris flower samples (can be stylized representations or actual iris photos)
- ▶ Row 2: Reconstructions of the same samples using 2 PCs

Add text below each column showing “Original”, “2 PCs (96% var.)”. If using actual measurements, show the 4 feature values as bar charts for each sample, comparing original (blue bars) vs. reconstructed (orange bars). Include reconstruction error values.

With 2 components (96% variance), reconstruction is excellent. Original and reconstructed feature values are very close.

Interpretation of Components

What do the principal components actually represent?

The first principal component \mathbf{u}_1 has approximate loadings:

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The second principal component focuses on contrasts between sepal and petal dimensions.

Note: Component interpretation is data-dependent and requires domain knowledge!

Assumptions of PCA

PCA makes several implicit assumptions:

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- ▶ Principal components are linear combinations of original features
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3. Orthogonality

- ▶ Components must be uncorrelated (orthogonal)
- ▶ Sometimes natural structure isn’t orthogonal

When PCA Struggles

[IMAGE PLACEHOLDER]

Description: Two examples where PCA fails:

- ▶ Left panel: “Swiss roll” - a 2D manifold embedded in 3D that looks like a rolled-up sheet. Show the 3D structure with points colored by their position along the roll. The structure is nonlinear—unrolling it requires nonlinear methods. Show that PCA would capture the major linear trends but miss the rolled structure.
- ▶ Right panel: Two classes in 2D that are separable but where the separation is in the low-variance direction. For example, two elongated parallel clusters with large variance along their length but small variance between them. PCA’s first component would go along the clusters (useless for classification), while the discriminative direction is the second component (low variance).

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Solutions:

- ▶ Robust PCA variants (using robust covariance estimators)
- ▶ Outlier detection and removal before PCA
- ▶ Regularization methods

Connections to Other Methods

PCA is related to many other techniques:

Probabilistic PCA (Bishop §12.2)

- ▶ Probabilistic latent variable model
- ▶ Provides principled handling of missing data
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Kernel PCA

- ▶ Nonlinear generalization using kernel trick
- ▶ Can capture curved manifold structure

More Extensions

Autoencoders

- ▶ Neural network approach to dimensionality reduction
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Supervised alternatives

- ▶ Linear Discriminant Analysis (LDA): uses class labels
- ▶ Partial Least Squares: considers both features and targets
- ▶ Better when goal is prediction, not just description

Practical Pitfalls

Common mistakes when applying PCA:

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3. Wrong number of components

- ▶ Too few: lose important information
- ▶ Too many: keep noise, computational waste
- ▶ Always validate on downstream task when possible

Implementation Options

Three main approaches in Python:

1. NumPy (manual implementation)

- ▶ Full control, educational value
- ▶ Use `np.cov()` and `np.linalg.eig()`
- ▶ Good for understanding, tedious for production

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3. SciPy

- ▶ Lower-level linear algebra routines
- ▶ `scipy.linalg` for SVD-based approach
- ▶ More flexible but requires more code

Scikit-learn Example: Basic Usage

```
1 import numpy as np
2 from sklearn.decomposition import PCA
3 from sklearn.preprocessing import StandardScaler
4
5 # Load your data (N samples, D features)
6 X = np.loadtxt('data.csv', delimiter=',')
7
8 # IMPORTANT: Standardize features to unit variance
9 scaler = StandardScaler()
10 X_scaled = scaler.fit_transform(X)
11
12 # Fit PCA - reduce to 2 dimensions
13 pca = PCA(n_components=2)
14 X_reduced = pca.fit_transform(X_scaled)
15
16 # X_reduced now has shape (N, 2)
17 print(f"Reduced data shape: {X_reduced.shape}")
```

Analyzing Results

```
1 # Variance explained by each component
2 print("Variance explained ratio:")
3 print(pca.explained_variance_ratio_)
4 # Output: [0.729, 0.229] (example values)
5
6 # Cumulative variance
7 print("Cumulative variance:")
8 print(pca.explained_variance_ratio_.cumsum())
9 # Output: [0.729, 0.958]
10
11 # Access principal components (shape: n_components x n_features
12 # )
12 components = pca.components_
13 print(f"PC1 loadings: {components[0]}")
14
15 # Reconstruct original data (with information loss)
16 X_reconstructed = pca.inverse_transform(X_reduced)
17
18 # Compute reconstruction error
```

Choosing Number of Components

```
1 # Strategy 1: Fit with all components first, then decide
2 pca_full = PCA()
3 pca_full.fit(X_scaled)
4
5 # Plot scree plot
6 import matplotlib.pyplot as plt
7 plt.figure(figsize=(8, 5))
8 plt.plot(range(1, len(pca_full.explained_variance_) + 1),
9         pca_full.explained_variance_, 'bo-')
10 plt.xlabel('Principal Component')
11 plt.ylabel('Eigenvalue')
12 plt.title('Scree Plot')
13 plt.show()
14
15 # Strategy 2: Specify variance threshold automatically
16 pca_auto = PCA(n_components=0.95) # Keep 95% variance
17 pca_auto.fit(X_scaled)
18 print(f"Number of components selected: {pca_auto.n_components_}")
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What is PCA?

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- ▶ Projects data onto lower-dimensional subspace
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Key assumptions:

- ▶ Linearity of projections
- ▶ Variance indicates importance
- ▶ Orthogonality of components

When to Use PCA

Excellent for:

- ▶ Exploratory data analysis
- ▶ Visualization of high-dimensional data (reduce to 2D/3D)
- ▶ Preprocessing for supervised learning (reduce features, speed up training)
- ▶ Noise reduction (keep top components, discard noisy ones)
- ▶ Compression (store low-dimensional representation)
- ▶ When data is approximately Gaussian with linear structure

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Consider alternatives when:

- ▶ Data lies on nonlinear manifold → Kernel PCA, t-SNE, UMAP
- ▶ Discriminative task with labels → LDA, supervised methods
- ▶ Need interpretable/sparse components → Sparse PCA, Factor Analysis
- ▶ Outliers present → Robust PCA

Looking Ahead

Topics we haven't covered:

Probabilistic PCA (Bishop §12.2)

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- ▶ Principled treatment of noise and missing data
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Advanced topics:

- ▶ Kernel PCA for nonlinear dimensionality reduction
- ▶ Factor Analysis and its relationship to PCA
- ▶ Modern deep learning approaches (variational autoencoders)
- ▶ Manifold learning methods (Isomap, LLE, t-SNE, UMAP)

Looking Ahead

Topics we haven't covered:

Probabilistic PCA (Bishop §12.2)

- ▶ Latent variable model: $\mathbf{x} = \mathbf{W}\mathbf{z} + \boldsymbol{\mu} + \boldsymbol{\epsilon}$
- ▶ Principled treatment of noise and missing data
- ▶ Enables Bayesian inference, model selection

Advanced topics:

- ▶ Kernel PCA for nonlinear dimensionality reduction
- ▶ Factor Analysis and its relationship to PCA
- ▶ Modern deep learning approaches (variational autoencoders)
- ▶ Manifold learning methods (Isomap, LLE, t-SNE, UMAP)

For supervised learning:

- ▶ Linear Discriminant Analysis (uses class labels)
- ▶ Canonical Correlation Analysis (two sets of variables)

References and Further Reading

Primary source:

- ▶ Bishop, C.M. (2006). *Pattern Recognition and Machine Learning*. Chapter 12 (§12.1)

Additional resources:

- ▶ Jolliffe, I.T. (2002). *Principal Component Analysis*. Springer. [Comprehensive treatment]
- ▶ Shlens, J. (2014). "A Tutorial on Principal Component Analysis." arXiv:1404.1100 [Accessible tutorial]
- ▶ James et al. (2013). *An Introduction to Statistical Learning*. Chapter 10.2 [Practical perspective]

Software documentation:

- ▶ Scikit-learn PCA guide:
scikit-learn.org/stable/modules/decomposition.html

Questions?