Lecture 18 - Class Activity: NMDS and PERMANOVA

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# Lecture 18: Non-metric Multidimensional Scaling (NMDS) and PERMANOVA

## What is NMDS?

NMDS (Non-metric Multidimensional Scaling) is an ordination technique that: - Visualizes dissimilarity between objects in reduced dimensions - Preserves rank order of distances, not exact distances - Works well with non-linear ecological relationships - Makes few assumptions about data structure

## When to Use NMDS

Use NMDS when you have: - **Community data**: Species abundance or presence/absence matrices - **Non-linear relationships**: When PCA assumptions are violated - **Complex ecological gradients**: Multiple environmental factors affecting communities

## Key Concepts of NMDS

1. **Dissimilarity matrices** instead of covariance
2. **Stress values** measure goodness of fit (<0.2 is acceptable)
3. **Iterative algorithm** to find optimal configuration
4. **No eigenvalues** - axes have no inherent meaning
5. **Rank-based** - preserves order, not exact distances

|  |  |
| --- | --- |
|  | **Critical First Step**Always check your stress value! Stress < 0.1 is excellent, 0.1-0.2 is good, > 0.2 is poor representation. |

# Part 1: Data Preparation and Exploration

## Load and Prepare Data

We’ll use the iris dataset for this analysis, treating it as if the measurements represent abundances of different “species” at different “sites”.

# Load iris data
iris\_df <- read\_csv("data/iris.csv") %>%
 clean\_names()

Rows: 150 Columns: 5
── Column specification ────────────────────────────────────────────────────────
Delimiter: ","
chr (1): species
dbl (4): sepal\_length, sepal\_width, petal\_length, petal\_width

ℹ Use `spec()` to retrieve the full column specification for this data.
ℹ Specify the column types or set `show\_col\_types = FALSE` to quiet this message.

# View data structure
head(iris\_df)

# A tibble: 6 × 5
 sepal\_length sepal\_width petal\_length petal\_width species
 <dbl> <dbl> <dbl> <dbl> <chr>
1 5.1 3.5 1.4 0.2 setosa
2 4.9 3 1.4 0.2 setosa
3 4.7 3.2 1.3 0.2 setosa
4 4.6 3.1 1.5 0.2 setosa
5 5 3.6 1.4 0.2 setosa
6 5.4 3.9 1.7 0.4 setosa

# For NMDS, we need just the numeric columns (our "species" data)
# We'll keep species as our grouping variable
iris\_species\_df <- iris\_df %>%
 dplyr::select(species)

iris\_numeric\_df <- iris\_df %>%
 dplyr::select(-species)

# Check the structure
str(iris\_numeric\_df)

tibble [150 × 4] (S3: tbl\_df/tbl/data.frame)
 $ sepal\_length: num [1:150] 5.1 4.9 4.7 4.6 5 5.4 4.6 5 4.4 4.9 ...
 $ sepal\_width : num [1:150] 3.5 3 3.2 3.1 3.6 3.9 3.4 3.4 2.9 3.1 ...
 $ petal\_length: num [1:150] 1.4 1.4 1.3 1.5 1.4 1.7 1.4 1.5 1.4 1.5 ...
 $ petal\_width : num [1:150] 0.2 0.2 0.2 0.2 0.2 0.4 0.3 0.2 0.2 0.1 ...

## Visualize the Data

# Create a pairs plot to see relationships
iris\_pairs\_plot <- iris\_df %>%
 dplyr::select(-species) %>%
 pairs(main = "Iris Measurements Relationships")



# Part 2: Running NMDS

## Step 1: Calculate Distance Matrix

# Calculate Bray-Curtis distance (common for ecological data)
# For iris data, we'll use Euclidean distance since these are measurements
iris\_dist <- dist(iris\_numeric\_df, method = "euclidean")

# Check the first few distances
iris\_dist[1:5]

[1] 0.5385165 0.5099020 0.6480741 0.1414214 0.6164414

## Step 2: Run NMDS

# Run NMDS with 2 dimensions
set.seed(123) # For reproducibility
iris\_nmds\_model <- metaMDS(iris\_numeric\_df,
 distance = "euclidean",
 k = 2, # Number of dimensions
 trymax = 100) # Maximum iterations

Run 0 stress 0.02525035
Run 1 stress 0.04045544
Run 2 stress 0.03566855
Run 3 stress 0.0268287
Run 4 stress 0.03695952
Run 5 stress 0.03104439
Run 6 stress 0.02682881
Run 7 stress 0.03962709
Run 8 stress 0.03210416
Run 9 stress 0.04325555
Run 10 stress 0.02525031
... New best solution
... Procrustes: rmse 1.741237e-05 max resid 7.129696e-05
... Similar to previous best
Run 11 stress 0.02525038
... Procrustes: rmse 1.217807e-05 max resid 8.876481e-05
... Similar to previous best
Run 12 stress 0.04000578
Run 13 stress 0.03552087
Run 14 stress 0.0309981
Run 15 stress 0.04078306
Run 16 stress 0.04008785
Run 17 stress 0.03197035
Run 18 stress 0.02525033
... Procrustes: rmse 5.481604e-06 max resid 2.712782e-05
... Similar to previous best
Run 19 stress 0.02682875
Run 20 stress 0.03270101
\*\*\* Best solution repeated 3 times

# Check the results
iris\_nmds\_model

Call:
metaMDS(comm = iris\_numeric\_df, distance = "euclidean", k = 2, trymax = 100)

global Multidimensional Scaling using monoMDS

Data: iris\_numeric\_df
Distance: euclidean

Dimensions: 2
Stress: 0.02525031
Stress type 1, weak ties
Best solution was repeated 3 times in 20 tries
The best solution was from try 10 (random start)
Scaling: centring, PC rotation
Species: expanded scores based on 'iris\_numeric\_df'

**Interpretation**: - Stress value: 0.025 - This is excellent representation

## Step 3: Extract NMDS Scores

# Extract NMDS scores for plotting
nmds\_scores\_df <- as.data.frame(iris\_nmds\_model$points) %>%
 rename(nmds1 = MDS1, nmds2 = MDS2) %>%
 bind\_cols(iris\_species\_df)

# View the scores
head(nmds\_scores\_df)

 nmds1 nmds2 species
1 -2.687311 0.2758924 setosa
2 -2.717262 -0.1705943 setosa
3 -2.882071 -0.1017541 setosa
4 -2.746720 -0.2603709 setosa
5 -2.730486 0.2902002 setosa
6 -2.311408 0.6625723 setosa

## Step 4: Create NMDS Plot

# Basic NMDS plot
nmds\_basic\_plot <- ggplot(nmds\_scores\_df, aes(x = nmds1, y = nmds2, color = species)) +
 geom\_point(size = 3, alpha = 0.8) +
 labs(title = "NMDS Ordination of Iris Data",
 subtitle = paste("Stress =", round(iris\_nmds\_model$stress, 3)),
 x = "NMDS1",
 y = "NMDS2",
 color = "Species") +
 theme\_minimal()

nmds\_basic\_plot



## Step 5: Add Confidence Ellipses

# NMDS plot with ellipses
nmds\_ellipse\_plot <- ggplot(nmds\_scores\_df, aes(x = nmds1, y = nmds2, color = species)) +
 geom\_point(size = 3, alpha = 0.8) +
 stat\_ellipse(level = 0.95, size = 1) +
 labs(title = "NMDS Ordination with 95% Confidence Ellipses",
 subtitle = paste("Stress =", round(iris\_nmds\_model$stress, 3)),
 x = "NMDS1",
 y = "NMDS2",
 color = "Species") +
 theme\_minimal()

Warning: Using `size` aesthetic for lines was deprecated in ggplot2 3.4.0.
ℹ Please use `linewidth` instead.

nmds\_ellipse\_plot



## Step 6: Stress Plot (Shepard Diagram)

# Create stress plot to evaluate fit
stressplot(iris\_nmds\_model, main = "Shepard Diagram: Ordination vs Original Distances")



# Part 3: PERMANOVA Analysis

## What is PERMANOVA?

PERMANOVA (Permutational Multivariate Analysis of Variance) tests whether groups have different multivariate centroids using permutation tests.

## Step 1: Run PERMANOVA

# Run PERMANOVA to test if species differ in multivariate space
set.seed(456)
iris\_permanova\_model <- adonis2(iris\_numeric\_df ~ species,
 data = iris\_df,
 method = "euclidean",
 permutations = 999)

# View results
iris\_permanova\_model

Permutation test for adonis under reduced model
Permutation: free
Number of permutations: 999

adonis2(formula = iris\_numeric\_df ~ species, data = iris\_df, permutations = 999, method = "euclidean")
 Df SumOfSqs R2 F Pr(>F)
Model 2 592.07 0.86894 487.33 0.001 \*\*\*
Residual 147 89.30 0.13106
Total 149 681.37 1.00000
---
Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

**Interpretation**: - F-statistic: 487.33 - R² (variance explained): 0.869 - p-value: 0.001

## Step 2: Check Homogeneity of Dispersions

Before interpreting PERMANOVA, we need to check if groups have similar multivariate spread.

# Test homogeneity of multivariate dispersions
iris\_dist\_full <- dist(iris\_numeric\_df)
dispersion\_model <- betadisper(iris\_dist\_full, iris\_df$species)

# Test for differences in dispersion
dispersion\_test <- anova(dispersion\_model)
dispersion\_test

Analysis of Variance Table

Response: Distances
 Df Sum Sq Mean Sq F value Pr(>F)
Groups 2 2.9092 1.45458 10.748 4.4e-05 \*\*\*
Residuals 147 19.8941 0.13533
---
Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

## Step 3: Visualize Dispersions

# Plot dispersions
plot(dispersion\_model, main = "Multivariate Dispersion by Species")



## Step 4: Pairwise PERMANOVA

# Function for pairwise PERMANOVA comparisons
pairwise\_permanova <- function(data\_matrix, groups, distance\_method = "euclidean") {
 # Get unique group combinations
 group\_levels <- unique(groups)
 comparisons <- combn(group\_levels, 2)

 # Initialize results
 results\_df <- data.frame(
 group1 = character(),
 group2 = character(),
 f\_statistic = numeric(),
 r\_squared = numeric(),
 p\_value = numeric()
 )

 # Run pairwise comparisons
 for(i in 1:ncol(comparisons)) {
 # Subset data
 group1 <- comparisons[1, i]
 group2 <- comparisons[2, i]
 subset\_indices <- which(groups %in% c(group1, group2))

 subset\_data <- data\_matrix[subset\_indices, ]
 subset\_groups <- groups[subset\_indices]

 # Run PERMANOVA
 temp\_result <- adonis2(subset\_data ~ subset\_groups,
 method = distance\_method,
 permutations = 999)

 # Store results
 results\_df <- rbind(results\_df, data.frame(
 group1 = group1,
 group2 = group2,
 f\_statistic = temp\_result$F[1],
 r\_squared = temp\_result$R2[1],
 p\_value = temp\_result$"Pr(>F)"[1]
 ))
 }

 # Adjust p-values for multiple comparisons
 results\_df$p\_adjusted <- p.adjust(results\_df$p\_value, method = "bonferroni")

 return(results\_df)
}

# Run pairwise comparisons
pairwise\_results\_df <- pairwise\_permanova(iris\_numeric\_df, iris\_df$species)
pairwise\_results\_df

 group1 group2 f\_statistic r\_squared p\_value p\_adjusted
1 setosa versicolor 551.0039 0.8489994 0.001 0.003
2 setosa virginica 943.7992 0.9059320 0.001 0.003
3 versicolor virginica 86.7697 0.4696100 0.001 0.003

# Part 4: ANOSIM Analysis

## What is ANOSIM?

ANOSIM (Analysis of Similarities) tests whether there is a significant difference between groups using rank dissimilarities.

## Step 1: Run ANOSIM

# Run ANOSIM
set.seed(789)
iris\_anosim\_model <- anosim(iris\_dist\_full, iris\_df$species, permutations = 999)

# View results
iris\_anosim\_model

Call:
anosim(x = iris\_dist\_full, grouping = iris\_df$species, permutations = 999)
Dissimilarity: euclidean

ANOSIM statistic R: 0.8794
 Significance: 0.001

Permutation: free
Number of permutations: 999

**Interpretation**: - R statistic: 0.879 - p-value: 0.001 - R close to 1 indicates strong separation between groups

## Step 2: Plot ANOSIM Results

# Plot ANOSIM results
plot(iris\_anosim\_model, main = "ANOSIM Results: Distribution of Permuted R Statistics")



# Part 5: Environmental Fitting (Optional)

If we had environmental variables, we could fit them to the ordination.

# For demonstration, let's use petal\_length as an "environmental" variable
env\_data\_df <- data.frame(petal\_length = iris\_df$petal\_length)

# Fit environmental vector
env\_fit\_model <- envfit(iris\_nmds\_model, env\_data\_df, permutations = 999)
env\_fit\_model

\*\*\*VECTORS

 NMDS1 NMDS2 r2 Pr(>r)
petal\_length 0.98205 -0.18862 0.9979 0.001 \*\*\*
---
Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1
Permutation: free
Number of permutations: 999

## Visualize Environmental Vectors

# Extract vector coordinates
env\_coords\_df <- as.data.frame(env\_fit\_model$vectors$arrows \* 2) # Scale for visibility
env\_coords\_df$variable <- rownames(env\_coords\_df)

# NMDS plot with environmental vector
nmds\_env\_plot <- ggplot(nmds\_scores\_df, aes(x = nmds1, y = nmds2, color = species)) +
 geom\_point(size = 3, alpha = 0.8) +
 stat\_ellipse(level = 0.95, size = 1, alpha = 0.3) +
 geom\_segment(data = env\_coords\_df,
 aes(x = 0, y = 0, xend = NMDS1, yend = NMDS2),
 arrow = arrow(length = unit(0.3, "cm")),
 color = "black", size = 1) +
 geom\_text(data = env\_coords\_df,
 aes(x = NMDS1 \* 1.1, y = NMDS2 \* 1.1, label = variable),
 color = "black", size = 4) +
 labs(title = "NMDS with Environmental Vector",
 subtitle = paste("Stress =", round(iris\_nmds\_model$stress, 3)),
 x = "NMDS1",
 y = "NMDS2",
 color = "Species") +
 theme\_minimal()

nmds\_env\_plot



# Summary Checklist for NMDS and PERMANOVA

|  |  |
| --- | --- |
|  | **Analysis Checklist**1. **Prepare your data** - ensure numeric matrix format
2. **Choose appropriate distance measure**
	* Bray-Curtis for abundance data
	* Euclidean for measurement data
3. **Run NMDS** with sufficient iterations
4. **Check stress value** - must be < 0.2
5. **Create ordination plots** with groups identified
6. **Test homogeneity of dispersions** before PERMANOVA
7. **Run PERMANOVA** to test group differences
8. **Consider ANOSIM** as complementary test
9. **Fit environmental variables** if available
 |

## Key Points to Remember

* **NMDS preserves rank order** of distances, not exact values
* **Stress < 0.2** is acceptable, < 0.1 is excellent
* **PERMANOVA tests centroids**, ANOSIM tests overlap
* **Check dispersion homogeneity** - violated assumption affects interpretation
* **Multiple comparisons** require p-value adjustment
* **Axes have no inherent meaning** in NMDS (unlike PCA)
* **Use appropriate distance measures** for your data type

|  |  |
| --- | --- |
|  | **Key Takeaways from NMDS/PERMANOVA Analysis**1. **NMDS is flexible** - works with any distance measure and makes few assumptions
2. **Stress indicates fit quality** - always report and check this value
3. **PERMANOVA is powerful** but assumes homogeneous dispersions
4. **ANOSIM is complementary** - provides different perspective on group separation
5. **Visualization is crucial** - always plot your ordination results
6. **Environmental fitting** helps interpret ecological patterns
7. **Permutation tests** avoid distributional assumptions

Remember: NMDS is iterative and may find different solutions - always set a seed for reproducibility! |