

# Week 2: Statistical Analysis & Feature Importance

## Heart Failure Survival Analysis

MDST Project

Winter 2026

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# Quick Recap: Week 1 - Exploratory Data Analysis

## Dataset Overview:

- 299 heart failure patients
- 13 features (12 predictors + target)
- Target: DEATH\_EVENT (0/1)
- 68% survived, 32% died
- No missing values

## Feature Types:

- **Continuous:** age, ejection\_fraction, serum\_creatinine, serum\_sodium, time, platelets, CPK
- **Binary:** anaemia, diabetes, high\_blood\_pressure, sex, smoking

# Week 1 Observations (Visual)

**What we saw in the plots:**

**Differences between groups:**

- Died: **Higher** serum\_creatinine
- Died: **Lower** ejection\_fraction
- Died: **Shorter** follow-up time
- Died: **Older** patients

**No obvious differences:**

- Diabetes (similar in both)
- Sex (similar in both)
- Smoking (similar in both)

**This Week's Goal:** Use **statistical tests** to confirm these observations with numbers!

# From Visualization to Statistics

Week 1: EDA	Week 2: Statistics
"The boxplots look different"	"The difference is statistically significant ( $p < 0.05$ )"
"This feature seems important"	"Random Forest ranks it #1"
"These features look correlated"	"VIF = 1.3, no multicollinearity"

**Key Insight:** Visualization suggests, statistics confirms!

# A Summer Afternoon in Cambridge, 1920s

*“I can tell whether the milk was poured first, or the tea.”*

## The Scene:

A group of scientists and academics gathered for afternoon tea at Cambridge University. Among them was Dr. Muriel Bristol, an algologist (scientist who studies algae).

When offered a cup of tea, she politely declined – insisting that she preferred her tea prepared with **milk poured into the cup first**, before the tea.

*“I can taste the difference,”* she claimed.

The scientists laughed. **Surely that’s impossible!**

The order of mixing couldn’t possibly affect the taste... could it?

## “Prove it.”

The room fell silent. How could they test such a claim?

- If she guesses correctly once, is that proof? (*Could be luck...*)
- What if she gets 2 right? 3 right? (*Still could be luck...*)
- How many correct answers would **convince** them she has real ability?

**This is the fundamental question of statistics:**

*How do we distinguish real effects from random chance?*

# Enter: Ronald Fisher



One man at the table saw this as more than a parlor game.

**Ronald Fisher** – a young statistician working at the Rothamsted Experimental Station – realized this simple question about tea contained a profound scientific problem.

*"I can design an experiment to test this,"* he said.

What followed would revolutionize science forever.



# Who Was Ronald Fisher?

## The “Father of Modern Statistics”

- Born in London, 1890
- Studied mathematics at Cambridge
- Poor eyesight prevented WWI service
- Worked on agricultural experiments
- Revolutionized scientific methodology

## His Contributions:

- **P-value** – probability under null
- **Null hypothesis** – default assumption
- **ANOVA** – comparing groups
- **Maximum likelihood estimation**
- **Experimental design principles**

*“To call in the statistician after the experiment is done may be no more than asking him to perform a post-mortem examination.” – R.A. Fisher*

# Fisher's Elegant Experiment

## The Design:

- 1 Prepare **8 cups** of tea: 4 with milk first, 4 with tea first
- 2 Present them in **random order**
- 3 Tell the lady there are exactly 4 of each type
- 4 Ask her to identify which 4 cups had milk added first

## The Mathematics:

- Total ways to choose 4 from 8:  
$$\binom{8}{4} = \frac{8!}{4! \times 4!} = 70$$
- Only **1 way** to get all 4 correct
- $P(\text{perfect by chance}) = \frac{1}{70} = 1.4\%$

## Fisher's Reasoning:

If she gets all 4 correct, there's only a 1.4% chance she was guessing.

This is small enough to **reject** the idea that she's just lucky!

## Fisher formalized this into a framework:

- 1 **Null Hypothesis ( $H_0$ ):** The lady has no ability (just guessing)
- 2 **Alternative Hypothesis ( $H_1$ ):** The lady has real ability
- 3 **P-value:** Probability of the result if  $H_0$  is true
- 4 **Decision:** If p-value  $< 0.05$ , reject  $H_0$

**The Result:** Dr. Bristol correctly identified **all 8 cups!**

$$\text{P-value} = 1.4\% < 5\%$$

*She really could taste the difference.*

# From Tea Cups to Heart Failure

The exact same logic applies to our analysis:

Lady Tasting Tea	Heart Failure Analysis
Can she distinguish milk-first from tea-first?	Can serum creatinine distinguish survivors from non-survivors?
$H_0$ : She's guessing randomly	$H_0$ : No difference between groups
If $p < 0.05 \rightarrow$ real ability	If $p < 0.05 \rightarrow$ feature matters

## Fisher's Legacy:

- These methods are now used in **medicine**, **biology**, **psychology**, and **data science**
- Every " $p < 0.05$ " you see in research traces back to Fisher

# What is Correlation?

**Pearson Correlation** measures the **linear relationship** between two variables.

**Range:**  $-1$  to  $+1$

- $+1$ : Perfect positive (both increase together)
- $0$ : No linear relationship
- $-1$ : Perfect negative (one up, other down)

**Formula:**

$$r = \frac{\sum (x_i - \bar{x})(y_i - \bar{y})}{\sqrt{\sum (x_i - \bar{x})^2 \sum (y_i - \bar{y})^2}}$$

**Key Question:** Which features are correlated with death?

# Feature Correlations with DEATH\_EVENT

## Positive Correlations:

- serum\_creatinine: +0.29
- age: +0.25

Higher values → higher death risk

## Negative Correlations:

- time: -0.53
- ejection\_fraction: -0.27
- serum\_sodium: -0.20

Higher values → lower death risk

## Key Insight:

Follow-up time has the strongest correlation – but this is expected!

Patients who die have shorter follow-up periods.

**Warning:** 'time' is a “leaky” feature – we wouldn't know it in advance for prediction!

# T-Test: Comparing Group Means

**Question:** Is there a significant difference in feature values between survivors and non-survivors?

## How it works:

- 1 Calculate mean for each group
- 2 Measure the difference
- 3 Account for variability
- 4 Get a p-value

## Assumptions:

- Normal distribution
- Independent samples
- (Welch's t-test relaxes equal variance)

**Null Hypothesis:** No difference between group means ( $\mu_1 = \mu_2$ )

# Mann-Whitney U Test: Non-Parametric Alternative

**When to use:** Data is NOT normally distributed

## How it works:

- 1 Rank all values together
- 2 Sum ranks for each group
- 3 Compare rank sums

## Advantages:

- No normality assumption
- Robust to outliers
- Works with ordinal data

## Example:

Age values: [55, 60, 65, 70, 75]

Ranks: [1, 2, 3, 4, 5]

If survivors have mostly low ranks and non-survivors have high ranks, there's a significant difference.



## Significant Features ( $p < 0.05$ )

Feature	T-test p-value	Mann-Whitney p-value	Significant?
time	$2.3 \times 10^{-22}$	$6.9 \times 10^{-21}$	Yes
ejection_fraction	$9.6 \times 10^{-6}$	$7.4 \times 10^{-7}$	Yes
serum_creatinine	$6.4 \times 10^{-5}$	$1.6 \times 10^{-10}$	Yes
age	$4.7 \times 10^{-5}$	$1.7 \times 10^{-4}$	Yes
serum_sodium	$1.9 \times 10^{-3}$	$2.9 \times 10^{-4}$	Yes
diabetes	0.97	0.97	No
sex	0.94	0.94	No
smoking	0.83	0.83	No

# The Problem with Multiple Testing

**Scenario:** Testing 12 features at  $\alpha = 0.05$

## The Math:

- Each test: 5% chance of false positive
- 12 tests: Expected false positives  
 $= 12 \times 0.05 = 0.6$
- Probability of at least one false positive:  
 $1 - (0.95)^{12} = 46\%$

## Real-World Problem:

- Publish 20 studies
- 1 will show “significant” result by chance
- This is why many studies don’t replicate!

**Solution:** Adjust p-values to control the False Discovery Rate (FDR)

# Benjamini-Hochberg (FDR) Correction

**Goal:** Control the expected proportion of false discoveries

**Procedure:**

- 1 Rank p-values from smallest to largest:  $p_{(1)} \leq p_{(2)} \leq \dots \leq p_{(m)}$
- 2 For each p-value, calculate the adjusted value:

$$p_{adj} = p \times \frac{m}{\text{rank}}$$

- 3 Compare adjusted p-values to  $\alpha$

## Why FDR over Bonferroni?

- Bonferroni: Very conservative ( $\alpha/m$ ) – may miss real effects
- FDR: Allows some false positives, but controls the rate

# After FDR Correction

## Still significant ( $\text{FDR} < 0.05$ ):

- time, ejection\_fraction, age, serum\_creatinine, serum\_sodium

## Not significant after correction:

- high\_blood\_pressure, anaemia, diabetes, platelets, sex, smoking, creatinine\_phosphokinase

**Conclusion:** Our 5 significant features remain significant even after accounting for multiple testing!

# Statistical Tests vs. Feature Importance

Statistical Tests	Feature Importance
“Is there a difference between groups?”	“How useful for prediction?”
Tests one feature at a time	Considers all features together
Measures statistical significance	Measures predictive power
Needs assumptions (normality, etc.)	Often model-based

**Key Insight:** A feature can be statistically significant but not useful for prediction (and vice versa). We need BOTH perspectives!

# What is Random Forest?

**Random Forest** = A collection of many decision trees that “vote” together

## Analogy:

Like asking 100 doctors for their opinion:

- Each doctor (tree) looks at different aspects
- They all vote on the outcome
- Majority vote wins

## Why Random Forest?

- Automatically learns useful features
- Captures non-linear relationships
- Robust and widely used
- Provides feature importance for free!

# Gini Importance (Mean Decrease in Impurity)

## How it works:

- 1 At each split, the tree asks: “Which feature best separates survived vs. died?”
- 2 Features that create better splits are used more often
- 3 Importance = how much each feature reduces “impurity” (mixing of classes)

## Simple Example:

- If `ejection_fraction` < 30 perfectly separates patients → HIGH importance
- If `smoking` doesn't help separate groups → LOW importance

**Limitation:** Can be biased toward features with many unique values

# Permutation Importance: A Better Alternative

**Problem with Gini:** Biased toward features with many values

## Permutation Importance Solution:

- 1 Train model and measure accuracy
- 2 Randomly shuffle one feature's values
- 3 Measure how much accuracy drops
- 4 Bigger drop = more important feature

**Intuition:** If shuffling a feature hurts the model a lot, that feature was important!

## Advantages:

- Unbiased
- Works with any model
- Evaluated on test data (more reliable)



# Feature Importance Results

## Top 5 Most Important Features:

- 1 **time** – Follow-up period (but “leaky”!)
- 2 **serum\_creatinine** – Kidney function indicator
- 3 **ejection\_fraction** – Heart pumping efficiency
- 4 **age** – Patient age
- 5 **serum\_sodium** – Electrolyte balance

## Clinical Interpretation:

- **Serum creatinine:** High levels indicate kidney dysfunction
- **Ejection fraction:** Low values mean heart isn't pumping efficiently
- These match the original research paper findings!

# Why Does Feature Variance Matter?

**Core Idea:** A feature that doesn't vary can't help distinguish groups!

**Example:**

- If ALL patients have diabetes = 1
- This feature tells us nothing
- Can't distinguish survivors from non-survivors

**Variance Formula:**

$$\text{Var}(X) = \frac{1}{n} \sum_{i=1}^n (x_i - \bar{x})^2$$

Measures spread around the mean

**Rule:** Remove features with very low variance (they carry no information)

# Coefficient of Variation (CV)

**Problem:** Raw variance depends on scale

- Platelets: variance =  $9.5 \times 10^9$  (large numbers!)
- Anaemia: variance = 0.25 (binary 0/1)
- Can't compare directly!

**Solution: Coefficient of Variation**

$$CV = \frac{\text{Standard Deviation}}{\text{Mean}} = \frac{\sigma}{\mu}$$

- Scale-free measure of relative spread
- $CV > 1$ : High variability relative to mean
- $CV < 1$ : Low variability relative to mean

# What is Multicollinearity?

**Definition:** When two or more features are highly correlated

## Simple Example:

- Height in cm
- Height in inches
- Same information!

## In Our Data:

- sex and smoking:  $r = 0.45$
- (Men smoke more in this dataset)

## Why is it a problem?

- Redundant information
- Hard to interpret importance
- Unstable regression coefficients
- Wastes model capacity

# Variance Inflation Factor (VIF)

**VIF** measures how much a feature's variance is “inflated” by correlation with others

## How it works:

- 1 For feature  $X_j$ , predict it using ALL other features
- 2 Calculate  $R_j^2$  (how well others predict  $X_j$ )
- 3 VIF formula:  $VIF_j = \frac{1}{1-R_j^2}$

## Intuition:

- If  $X_j$  predicted perfectly by others ( $R^2 = 1$ ):  $VIF \rightarrow \infty$
- If  $X_j$  is independent ( $R^2 = 0$ ):  $VIF = 1$

VIF Value	Interpretation & Action
$VIF = 1$	No correlation – no action needed
$1 < VIF < 5$	Moderate – usually acceptable
$5 \leq VIF < 10$	High – investigate further
$VIF \geq 10$	Severe – consider removing feature

## What to do with high VIF?

- Remove one of the correlated features
- Combine features (e.g., PCA)
- Use regularization (Ridge, Lasso)

# VIF in Python

```
from statsmodels.stats.outliers_influence
    import variance_inflation_factor
from sklearn.preprocessing import StandardScaler

# Scale features for numerical stability
X_scaled = StandardScaler().fit_transform(X)

# Calculate VIF for each feature
vif_data = pd.DataFrame()
vif_data['Feature'] = X.columns
vif_data['VIF'] = [
    variance_inflation_factor(X_scaled, i)
    for i in range(X_scaled.shape[1])
]
vif_data.sort_values('VIF', ascending=False)
```

# VIF Results for Our Dataset

Feature	VIF	Status
sex	1.35	OK
smoking	1.32	OK
age	1.15	OK
time	1.14	OK
serum_sodium	1.13	OK
ejection_fraction	1.10	OK
serum_creatinine	1.08	OK

## Good news!

- All VIF values are close to 1
- No severe multicollinearity
- All features provide relatively independent information

**Note:** sex and smoking have slightly higher VIF due to their correlation (0.45)



# Variance vs. VIF: Key Differences

Variance	VIF
Measures spread of a <i>single</i> feature	Measures correlation <i>between</i> features
Low variance = feature doesn't vary much	High VIF = feature is redundant
Question: "Does this feature have different values?"	Question: "Is this feature's info already captured by others?"
Solution: Remove low-variance features	Solution: Remove one of correlated pair

# Key Takeaways

- 1 **Correlation** measures linear relationships between variables
- 2 **T-test** compares means (assumes normality)
- 3 **Mann-Whitney U** is non-parametric (no normality assumption)
- 4 **FDR Correction** controls false discovery rate in multiple testing
- 5 **Random Forest** provides feature importance scores
- 6 **Permutation Importance** is more reliable than Gini importance
- 7 **Variance/CV** tells us how much features vary
- 8 **VIF** detects multicollinearity (redundant features)

# Key Findings for Heart Failure Dataset

## Most Important Predictive Features:

- 1 time (but leaky!)
- 2 serum\_creatinine
- 3 ejection\_fraction
- 4 age
- 5 serum\_sodium

## Data Quality:

- No multicollinearity issues (all VIF  $< 5$ )
- All features have adequate variance
- Results match the original research paper!

# Next Week: Unsupervised Learning

- **PCA** (Principal Component Analysis)
  - Dimensionality reduction
  - Visualizing high-dimensional data
- **Clustering**
  - Finding natural groupings in data
  - K-means, hierarchical clustering

- 1 Write a function to return features with significant p-values given a threshold
- 2 Implement Mann-Whitney U test for all features
- 3 Compare feature rankings from t-test vs. Random Forest
- 4 Calculate VIF for all features and interpret results
- 5 Write a low-variance filter function

## Resources:

- Scipy Stats: <https://docs.scipy.org/doc/scipy/reference/stats.html>
- Statsmodels VIF: <https://www.statsmodels.org/>
- Scikit-learn: <https://scikit-learn.org/>