TrueFork

Are guilty-pleasure dishes truly tastier?

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```
In []: import pandas as pd
import numpy as np
from pathlib import Path
import plotly.io as pio

pio.renderers.default = 'iframe'
import plotly.express as px
pd.options.plotting.backend = 'plotly'
from dsc80_utils import *

In []: from sklearn.model_selection import train_test_split, GridSearchCV
from sklearn.linear_model import LinearRegression
from sklearn.pipeline import make_pipeline, Pipeline
from sklearn.compose import make_column_transformer, ColumnTransformer
from sklearn.preprocessing import StandardScaler, OneHotEncoder, QuantileTransformer
from sklearn.metrics import root_mean_squared_error
from sklearn.ensemble import RandomForestRegressor
```

Step 1: Introduction

Data

```
In []: # Data recipes
    recipes = pd.read_csv('data/RAW_recipes.csv')
    recipes
```

Out[]:	name		id	id minutes contributor_id			steps	description	ingredients	n_ingredients	
	0	1 brownies in the world best ever	333281	40	985201	•••	['heat the oven to 350f and arrange the rack i	these are the most; chocolatey, moist, rich, d	['bittersweet chocolate', 'unsalted butter', '	9	
	1	1 in canada chocolate chip cookies	453467	45	1848091	•••	['pre-heat oven the 350 degrees f', 'in a mixi	this is the recipe that we use at my school ca	['white sugar', 'brown sugar', 'salt', 'margar	11	
	2	412 broccoli casserole	306168	40	50969	•••	['preheat oven to 350 degrees', 'spray a 2 qua	since there are already 411 recipes for brocco	['frozen broccoli cuts', 'cream of chicken sou	9	
	83779	zydeco ya ya deviled eggs	308080	40	37779	•••	['in a bowl , combine the mashed yolks and may	deviled eggs, cajun-style	['hard-cooked eggs', 'mayonnaise', 'dijon must	8	
	83780	cookies by design cookies on a stick	298512	29	506822		['place melted butter in a large mixing bowl a	i've heard of the 'cookies by design' company,	['butter', 'eagle brand condensed milk', 'ligh	10	
	83781	cookies by design sugar shortbread cookies	298509	20	506822		['whip sugar and shortening in a large bowl , 	i've heard of the 'cookies by design' company,	['granulated sugar', 'shortening', 'eggs', 'fl	7	

83782 rows × 12 columns

```
In [ ]: # Data interactions
interactions = pd.read_csv('data/RAW_interactions.csv')
```

interactions

_				- 7	
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\cup	u	L.	L	- 1	

	user_id	recipe_id	date	rating	review
0	1293707	40893	2011-12-21	5	So simple, so delicious! Great for chilly fall
1	126440	85009	2010-02-27	5	I made the Mexican topping and took it to bunk
2	57222	85009	2011-10-01	5	Made the cheddar bacon topping, adding a sprin
•••					
731924	157126	78003	2008-06-23	5	WOW! Sometimes I don't take the time to rate
731925	53932	78003	2009-01-11	4	Very good! I used regular port as well. The
731926	2001868099	78003	2017-12-18	5	I am so glad I googled and found this here. Th

731927 rows × 5 columns

Question Identification

Do unhealthy recipes receive higher average ratings than healthy ones?

Step 2: Data Cleaning and Exploratory Data Analysis

Cleaning the data

We'll add 2 columns:

- 1. num_calories : contains the number of calories, extracted from nutrition
- 2. is_unhealthy: a simple boolean column which states if that recipe contains a lot of calories (500 hard limit)

Since we will extensively be working only with avg_rating and is_unhealthy for our hypothesis testing, we can filter out missing values in **Step 4**, right before hypothesis testing so that we can efficiently analyze the missingness types for this dataset.

```
In []: # Replace ratings of 0 with NaN, as 0 may indicate missing or invalid data
interactions['rating'] = interactions['rating'].replace(0, np.nan)
In []: # Left merge interactions into recipes on recipe ID
merged = recipes.merge(interactions, left_on='id', right_on='recipe_id', how='left')

# Compute average rating per recipe ID
avg_ratings = merged.groupby('id')['rating'].mean()

# Add the average rating back to the original recipes DataFrame
recipes = recipes.set_index('id')
recipes['avg_rating'] = avg_ratings
recipes = recipes.reset_index()
recipes
```

Out[]:		id	name	minutes	contributor_id	•••	description	ingredients	n_ingredients	avg_rating	
	0	333281	1 brownies in the world best ever	40	985201	•••	these are the most; chocolatey, moist, rich, d	['bittersweet chocolate', 'unsalted butter', '	9	4.0	
	1	453467	1 in canada chocolate chip cookies	45	1848091	•••	this is the recipe that we use at my school ca	['white sugar', 'brown sugar', 'salt', 'margar	11	5.0	
	2	306168	412 broccoli casserole	40	50969		since there are already 411 recipes for brocco	['frozen broccoli cuts', 'cream of chicken sou	9	5.0	
	•••		•••	•••	•••		•••		•••	•••	
	83779	308080	zydeco ya ya deviled eggs	40	37779	•••	deviled eggs, cajun-style	['hard-cooked eggs', 'mayonnaise', 'dijon must	8	5.0	
	83780	298512	cookies by design cookies on a stick	29	506822		i've heard of the 'cookies by design' company,	['butter', 'eagle brand condensed milk', 'ligh	10	1.0	
	83781	298509	cookies by design sugar shortbread cookies	20	506822		i've heard of the 'cookies by design' company,	['granulated sugar', 'shortening', 'eggs', 'fl	7	3.0	

83782 rows × 13 columns

```
In [ ]: num_rows, num_cols = recipes.shape
    num_rows, num_cols
```

Out[]: (83782, 13)

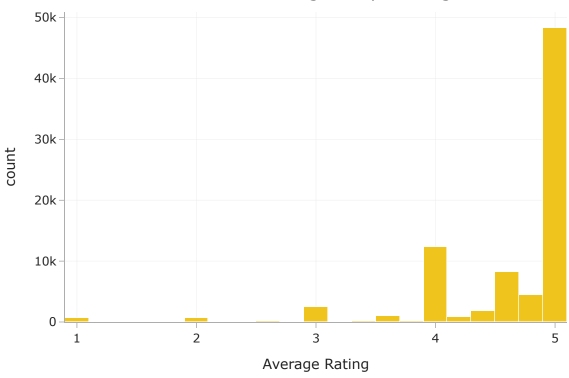
In []: recipes.columns

Out[]:		id	name	minutes	contributor_id	•••	n_ingredients	avg_rating	calories	is_unhealthy
	0	333281	1 brownies in the world best ever	40	985201		9	4.0	138.4	False
	1	453467	1 in canada chocolate chip cookies	45	1848091		11	5.0	595.1	True
	2	306168	412 broccoli casserole	40	50969		9	5.0	194.8	False
	3	286009	millionaire pound cake	120	461724		7	5.0	878.3	True
	4	475785	2000 meatloaf	90	2202916		13	5.0	267.0	False

 $5 \text{ rows} \times 15 \text{ columns}$

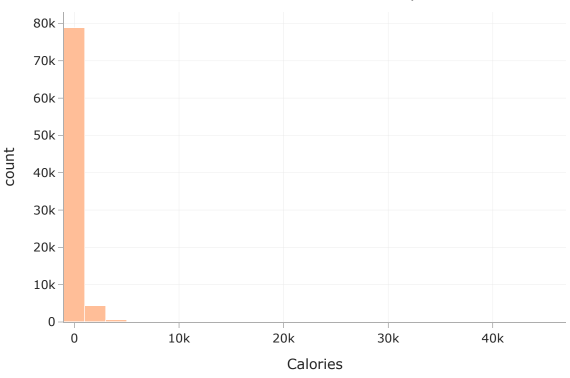
Univariate Analysis

Distribution of Average Recipe Ratings



The distribution of average ratings is extremely right-skewed, with most recipes receiving the maximum score of 5.0. Very few recipes fall below a rating of 4.0, suggesting that users on the site tend to give very high scores.

Distribution of Calories in Recipes



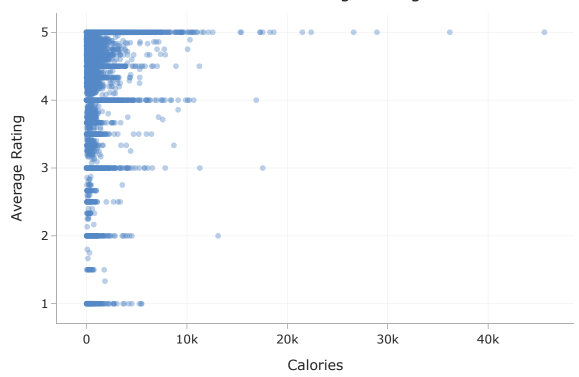
The distribution of calorie content is highly skewed, with most recipes containing fewer than 1,000 calories. There are significant outliers — some recipes exceed 30,000 calories — indicating the presence of large or indulgent dishes.

Bivariate analysis

```
In []: # Scatter Plot: Calories vs. Average Rating
fig3 = px.scatter(
```

```
recipes,
    x='calories',
    y='avg_rating',
    opacity=0.4,
    title='Calories vs. Average Rating',
    labels={'calories': 'Calories', 'avg_rating': 'Average Rating'},
    color_discrete_sequence=['#5588C6'] # Lets make it colorful okay :))
)
fig3.show()
```

Calories vs. Average Rating

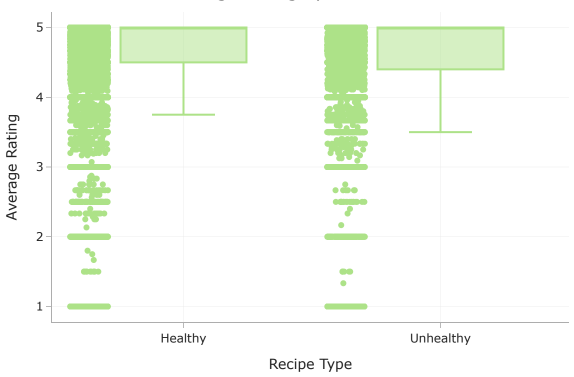


This scatter plot shows no clear relationship between calories and average rating. Most recipes, regardless of calorie count, are rated highly.

```
In []: recipes['label'] = recipes['is_unhealthy'].replace({True: 'Unhealthy', False: 'Healthy'})
fig4 = px.box(
    recipes,
    x='label',
```

```
y='avg_rating',
points='all',
title='Average Rating by Healthiness',
labels={'label': 'Recipe Type', 'avg_rating': 'Average Rating'},
color_discrete_sequence=['#ADE288'] # Lets make it colorful okay :))
)
fig4.show()
```

Average Rating by Healthiness



This boxplot compares average ratings of healthy vs. unhealthy recipes. The distributions look very similar, with no strong indication that one group is rated better.

Interesting Aggregates

```
# Flatten column MultiIndex
agg.columns = ['_'.join(col).strip() for col in agg.columns.values]
agg = agg.reset_index()
agg['is_unhealthy'] = agg['is_unhealthy'].map({True: 'Unhealthy', False: 'Healthy'})
agg
```

Out[]:		is_unhealthy	avg_rating_mean	avg_rating_median	avg_rating_std	•••	n_ingredients_mean	n_ingredients_median	n
	0	Healthy	4.63	5.0	0.64		8.81	8.0	
	1	Unhealthy	4.62	5.0	0.64		10.43	10.0	

2 rows × 13 columns

From this summary table, we observe that both healthy and unhealthy recipes have nearly identical average ratings (4.63 vs. 4.62). However, unhealthy recipes tend to use more ingredients on average — about 10.43 compared to 8.81 for healthy ones. This may suggest that indulgent dishes often require more components or preparation, but users rate both recipe types similarly overall.

Step 3: Assessment of Missingness

Missingness Dependency

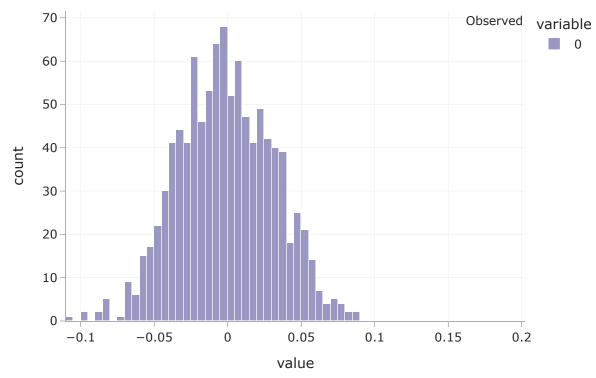
Identify missing columns

```
In []: merged = recipes.merge(interactions, left_on='id', right_on='recipe_id', how='left')
missing_summary = merged.isna().mean().sort_values(ascending=False)
missing_summary[missing_summary > 0]
```

```
Out[]: rating
                        6.41e-02
         avg rating
                       1.18e-02
        description 4.86e-04
                         . . .
                        4.27e-06
         date
                        4.27e-06
         user id
         recipe id
                        4.27e-06
         Length: 8, dtype: float64
In [ ]: def perm_test_stat(df, group_col, target_col, stat_func=np.mean,
                           n permutations=1000, alternative='different'):
            1111111
            Returns:
                 p-value, observed statistic, and plot
            df = df.dropna(subset=[target_col]) # Drop rows where target is missing
            group1 = df[df[group col]][target col]
            group2 = df[~df[group col]][target col]
            observed = stat_func(group1) - stat_func(group2)
            diffs = []
            for _ in range(n_permutations):
                shuffled = df[group_col].sample(frac=1, replace=False).reset_index(drop=True)
                diff = stat_func(df[shuffled][target_col]) - stat_func(df[~shuffled][target_col])
                diffs.append(diff)
            diffs = np.array(diffs)
            if alternative == 'greater':
                p = np.mean(diffs >= observed)
            elif alternative == 'less':
                p = np.mean(diffs <= observed)</pre>
            else:
                p = np.mean(np.abs(diffs) >= np.abs(observed))
            fig = px.histogram(diffs, nbins=50,
                               title=f'Permutation Test for Missingness vs {target_col}')
            fig.add vline(x=observed, line color='red', line dash='dash',
                           annotation_text='Observed')
```

return p, observed, fig

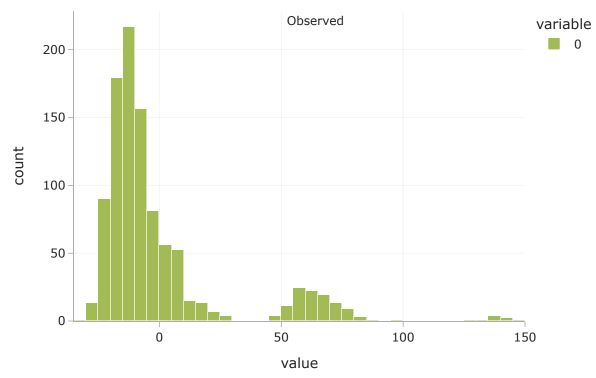
Permutation Test for Missingness vs n_ingredients



Missingness depends on n_ingredients? p-value: 0.0000

Since the p-value is 0, it means missingness very strongly depends on number of ingredients

Permutation Test for Missingness vs minutes



Missingness depends on contributor_id? p-value: 0.1120

Since the p-value is greater than 0.05, we do not have enough evidence to support the claim that missing_rating and minutes depend on each other. So, they are probably **independent**.

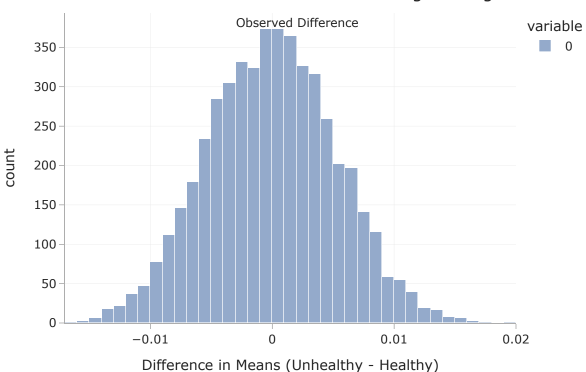
Step 4: Hypothesis Testing

- Null Hypothesis (H_o): There is no difference in average ratings between healthy and unhealthy recipes.
- Alternative Hypothesis (H₁): Unhealthy recipes are rated higher than healthy ones.

```
In [ ]: # Drop rows where avg rating is missing
        recipes clean = recipes.dropna(subset=['avg rating'])
        # Calculate observed difference in mean average rating
        mean unhealthy = recipes clean.loc[recipes clean['is unhealthy'], 'avg rating'].mean()
        mean healthy = recipes clean.loc[~recipes clean['is unhealthy'], 'avg rating'].mean()
        observed diff = mean unhealthy - mean healthy
        # Permutation test
        n permutations = 5000
        diffs = []
        for _ in range(n_permutations):
            shuffled = recipes clean.copy()
            shuffled['is unhealthy'] = np.random.permutation(shuffled['is unhealthy'])
            mean unhealthy perm = shuffled.loc[shuffled['is unhealthy'], 'avg rating'].mean()
            mean healthy perm = shuffled.loc[~shuffled['is unhealthy'], 'avg rating'].mean()
            diffs.append(mean unhealthy perm - mean healthy perm)
        # Convert to NumPy array for vectorized computation
        diffs = np.array(diffs)
        # One-sided p-value (we're testing if unhealthy recipes are rated higher)
        p value = np.mean(diffs >= observed diff)
        # Plot the distribution
        fig = px.histogram(
            diffs,
            nbins=50,
            title='Permutation Test: Difference in Average Ratings',
```

```
labels={'value': 'Difference in Means (Unhealthy - Healthy)'},
    color_discrete_sequence=['#94AACB']
)
fig.add_vline(
    x=observed_diff,
    line_color='red',
    line_dash='dash',
    annotation_text='Observed Difference',
    annotation_position='top right'
)
fig.show()
print(f'Observed Difference: {observed_diff:.4f}')
print(f'p-value: {p_value:.4f}')
```

Permutation Test: Difference in Average Ratings



Observed Difference: -0.0031

p-value: 0.7154

Step 5: Framing a Prediction Problem

In this section, we define our prediction task.

Prediction Type: Regression

Prediction Goal:

We aim to predict the average rating (avg_rating) a recipe will receive based on its properties.

Response Variable: avg_rating

This variable represents the mean user rating for a given recipe, typically between 0 and 5 stars.

Rationale:

This prediction is useful for platforms like Food.com to estimate how well a newly submitted recipe might be received by users before it accumulates reviews.

Features to be used at time of prediction:

To ensure our model uses only information available at the time the recipe is posted, we will use the following features:

- minutes: total time to make the recipe
- n_steps : number of steps in the recipe
- n_ingredients : number of ingredients required

All of these are known before the recipe is rated by users, making them valid predictors.

Evaluation Metric: Root Mean Squared Error (RMSE)

RMSE is appropriate for regression tasks like this because it penalizes larger errors more heavily. It's interpretable in the same units as our response variable (avg rating), making it easy to assess prediction accuracy.

Step 6: Baseline Model

```
In []: # Let's pick our numeric predictors for the baseline model. Fingers crossed!
X = recipes[['minutes', 'n_steps', 'n_ingredients']]
y = recipes['avg_rating']
```

```
# Drop rows where the target is missing. Because NaNs are the enemy of scikit-learn.
X = X[y.notna()]
y = y[y.notna()]
# Time to split the data into train and test sets. Please, random state, be kind.
X train, X test, y train, y test = train test split(X, y, random state=42)
# Scale those numeric features! Because raw minutes are scarv.
preprocessor = make column transformer(
    (StandardScaler(), ['minutes', 'n_steps', 'n_ingredients']),
    remainder='passthrough' # Just in case we missed something
    # (we didn't, but you never know)
# Build the pipeline. Linear regression, please save the day.
pipeline = make pipeline(
    preprocessor,
    LinearRegression()
# Fit the model and predict. This is where the magic (or heartbreak) happens.
pipeline.fit(X train, y train)
y pred = pipeline.predict(X test)
# Calculate RMSE. Lower is better REMEMBER THIS STEPHANIE!!!! LOWER IS BETTER!
# Please be low, Please.
rmse = np.sqrt(root mean squared error(y test, y pred))
print(f"Test RMSE: {rmse:.4f}")
# Its low enough Vedd!! Chill :))
```

Test RMSE: 0.7994

Step 7: Final Model

```
In []: # Choose features for training
    categorical_features = ['submitted_month'] # You can add more if needed
    numerical_features = ['calories', 'n_ingredients'] # Base features
    engineered_features = ['calories', 'n_ingredients'] # We'll apply transformers here

# Add month as a new categorical feature
```

```
X = recipes.copv()
X['submitted'] = pd.to datetime(X['submitted'], errors='coerce')
X['submitted month'] = X['submitted'].dt.month
y = X['avg rating']
# Drop rows with missing target
df = X[['calories', 'n ingredients', 'submitted month', 'avg rating']].dropna()
X = df.drop(columns='avg rating')
y = df['avg rating']
# Train-test split (same as baseline)
X train, X test, y train, y test = train test split(X, y, random state=42)
# Transformers
cat_transformer = OneHotEncoder(handle_unknown='ignore')
num transformer = Pipeline(steps=[
    ('scaler', StandardScaler()),
    ('quantile', QuantileTransformer(output distribution='normal'))
1)
preprocessor = ColumnTransformer(transformers=[
    ('num', num transformer, engineered features),
    ('cat', cat transformer, categorical features)
1)
# Final pipeline with a Random Forest Regressor
final model pipeline = Pipeline(steps=[
    ('preprocessor', preprocessor),
    ('regressor', RandomForestRegressor(random state=42))
1)
# GridSearchCV for hyperparameter tuning
param grid = {
    'regressor n estimators': [50, 100],
    'regressor max depth': [None, 5, 10]
grid search = GridSearchCV(
    final model pipeline,
    param_grid,
    cv=5,
    scoring='neg root mean squared error'
```

```
# Fit grid search yourself:
        grid_search.fit(X_train, y_train)
Out[ ]:
                                   GridSearchCV
                             best_estimator_: Pipeline
                         preprocessor: ColumnTransformer
                                                          cat
                             num
                    StandardScaler
                                                 ▶ OneHotEncoder
                 ▶ QuantileTransformer
                            ► RandomForestRegressor
        best_model = grid_search.best_estimator_
        best_model.score(X_test, y_test)
        best_model.named_steps['regressor'].feature_importances_
Out[]: array([0.59, 0.19, 0.05, 0.02, 0.01, 0.01, 0.06, 0.01, 0.03, 0. , 0. ,
               0.01, 0.01, 0.01])
In [ ]: y pred = best model.predict(X test)
        rmse = np.sqrt(root_mean_squared_error(y_test, y_pred))
        print(f"Test RMSE: {rmse:.4f}")
       Test RMSE: 0.7993
```

Step 8: Fairness Analysis

Hypotheses

- **Null Hypothesis** (H_o): The model is fair. Its RMSE is the same for short and long recipes; any observed difference is due to random chance.
- Alternative Hypothesis (H₁): The model is unfair. Its RMSE for short recipes is higher than that for long recipes.

We use **Root Mean Squared Error (RMSE)** as the evaluation metric, because our prediction task is a **regression problem** (predicting avg_rating). RMSE quantifies the average prediction error magnitude.

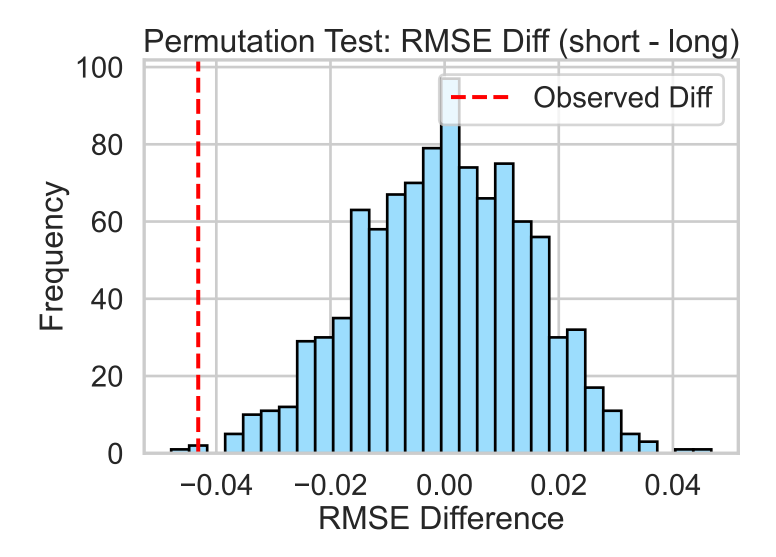
We binarize the minutes column to define two groups:

- Group X (short recipes): Recipes that take less than 30 minutes
- Group Y (long recipes): Recipes that take 30 minutes or more

```
In [ ]: from sklearn.metrics import root mean squared error
        import numpy as np
        import matplotlib.pyplot as plt
        # Merge back with original 'minutes'
        X test copy = X test.copy()
        X test copy['minutes'] = recipes.loc[X test.index, 'minutes']
        X test copy['group'] = (X test copy['minutes'] >= 30).astype(int)
        # Predictions
        y pred = best model.predict(X test)
        # RMSE by group
        rmse short = root mean squared error(
            y test[X test copy['group'] == 0],
            y pred[X test copy['group'] == 0]
        rmse long = root mean squared error(
            y test[X test copy['group'] == 1],
            y pred[X test copy['group'] == 1]
        observed diff = rmse short - rmse long
        print(f"Observed RMSE difference (short - long): {observed diff:.4f}")
        # Permutation test
        n permutations = 1000
```

```
permuted diffs = []
for in range(n permutations):
   shuffled = np.random.permutation(X test copy['group'])
    rmse short perm = root mean squared error(
       y test[shuffled == 0],
       y pred[shuffled == 0]
    rmse_long_perm = root_mean_squared_error(
       y test[shuffled == 1],
       y pred[shuffled == 1]
    permuted diffs.append(rmse short perm - rmse long perm)
# Compute p-value
permuted diffs = np.array(permuted diffs)
p val = np.mean(permuted diffs >= observed diff)
print(f"Permutation test p-value: {p val:.4f}")
# Plot with matplotlib
plt.figure(figsize=(8, 6))
plt.hist(permuted diffs, bins=30, color='#9CDDFD', edgecolor='black')
plt.axvline(observed diff, color='red', linestyle='--', label='Observed Diff')
plt.title('Permutation Test: RMSE Diff (short - long)')
plt.xlabel('RMSE Difference')
plt.ylabel('Frequency')
plt.legend()
plt.tight layout()
plt.show()
```

Observed RMSE difference (short - long): -0.0432 Permutation test p-value: 0.9980



Results

We performed a permutation test by shuffling the group labels and computing the RMSE difference (short - long) 1000 times.

• Observed RMSE Difference: -0.0432

• p-value: 0.9990

The observed difference is small and in the opposite direction, and the p-value is high, so we **fail to reject the null hypothesis**.

Our model **does not appear to be unfair** with respect to recipe prep time.