Homogenous Ensemble Learning in Highly Unbalanced Data

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Website Link: https://kevinbian107.github.io/ensemble-imbalanced-data/

```
In []: # for eda and modeling
   import pandas as pd
   import numpy as np
   pd.options.plotting.backend = 'plotly'
   from utils.dsc80_utils import *
   from itertools import chain
```

Step 1: Introduction

Predictive model detecting user preference using **textual features** in combnation with other **numerical features** is the key first step prior to building a recommander system or doing any other further analysis. The challenge that is addressed in this project is related to the high unbalance nature of the recipe data set that we are using.

```
In [ ]: interactions = pd.read_csv('food_data/RAW_interactions.csv')
    recipes = pd.read_csv('food_data/RAW_recipes.csv')
```

Step 2: Data Cleaning and Exploratory Data Analysis

Merging

Initial merging is needed for the two dataset to form 1 big data set

- 1. Left merge the recipes and interactions datasets together.
- 2. In the merged dataset, fill all ratings of 0 with np.nan. (Think about why this is a reasonable step, and include your justification in your website.)
- 3. Find the average rating per recipe, as a Series.
- 4. Add this Series containing the average rating per recipe back to the recipes dataset however you'd like (e.g., by merging). Use the resulting dataset for all of your analysis. (For the purposes of Project 4, the 'review' column in the interactions dataset doesn't have much use.)

Transformation

- 1. Some columns, like nutrition, contain values that look like lists, but are actually strings that look like lists. We turned the strings into actual columns for every unique value in those lists
- 2. Convert to list for steps , ingredients , and tags
- 3. Convert date and submitted to Timestamp object and rename as review_date and recipe_date
- 4. Convert Types
- 5. Drop same id (same with recipe_id)
- 6. Replace 'nan' with np.NaN

Type Logic

- 1. String: [name, contributor_id, user_id, recipe_id,]
 - quantitative or qualitative, but cannot perform mathamatical operations (quntitative discrete)

- name is the name of recipe
- contributor id is the author id of the recipe (shape=7157)
- recipe_id is the id of teh recipe (shape=25287)
 - id from the original dataframe also is the id of the recipe, dropped after merging
- user id is the id of the reviewer (shape=8402)
- 2. List: [tags, steps, description, ingredients, review]
 - qualitative, no mathamatical operation (qualitative discrete)
- 3. int : [n_steps, minutes, n_ingredients, rating]
 - quantitative mathamatical operations allowed (quantitative continuous)
- 4. float: [avg_rating, calories, total_fat sugar, sodium, protein, sat_fat, carbs]
 - quantitative mathamatical operations allowed (quantitative continuous)
- 5. Timestamp: [recipe_date, review_date]
 - quantitative mathamatical operations allowed (quantitative continuous)

Below are the full implementation of initial, which does the merge conversion, then transform, which carries out the neccessary transformation described above

```
In [ ]: def initial(df):
            '''Initial claeaning and megrging of two df, add average ratings'''
            # fill 0 with np.NaN
            df['rating'] = df['rating'].apply(lambda x: np.NaN if x==0 else x)
            # not unique recipe_id
            avg = df.groupby('recipe_id')[['rating']].mean().rename(columns={'rating':'avg_rating'})
            df = df.merge(avg, how='left', left_on='recipe_id',right_index=True)
            return df
        def transform df(df):
            '''Transforming nutrition to each of its own catagory,
            tags, steps, ingredients to list,
            submission date to timestamp object,
            convert types,
            and remove 'nan' to np.NaN'''
            # Convert nutrition to its own caatgory
            data = df['nutrition'].str.strip('[]').str.split(',').to_list()
            name = {0:'calories',1:'total_fat',2:'sugar',3:'sodium',4:'protein',5:'sat_fat',6:'carbs'}
            #zipped = data.apply(lambda x: list(zip(name, x)))
            new = pd.DataFrame(data).rename(columns=name)
            df = df.merge(new,how='inner',right_index=True, left_index=True)
            df = df.drop(columns=['nutrition'])
            # Convert to list
            def convert_to_list(text):
                return text.strip('[]').replace("'",'').split(', ')
            df['tags'] = df['tags'].apply(lambda x: convert to list(x))
            df['ingredients'] = df['ingredients'].apply(lambda x: convert to list(x))
            # it's correct, just some are long sentences, doesn't see "'", notice spelling
            df['steps'] = df['steps'].apply(lambda x: convert_to_list(x)) #some white space need to be h
            # submission date to time stamp object
            format ='%Y-%m-%d'
            df['submitted'] = pd.to_datetime(df['submitted'], format=format)
            df['date'] = pd.to_datetime(df['date'],format=format)
            # drop not needed & rename
            df = df.drop(columns=['id']).rename(columns={'submitted':'recipe_date','date':'review_date'}
            # Convert data type
            df[['calories','total_fat','sugar',
```

	name	minutes	contributor_id	recipe_date	 sodium	protein	sat_fat	carbs
0	1 brownies in the world best ever	40	985201	2008-10-27	 3.0	3.0	19.0	6.0
1	1 in canada chocolate chip cookies	45	1848091	2011-04-11	 22.0	13.0	51.0	26.0
2	412 broccoli casserole	40	50969	2008-05-30	 32.0	22.0	36.0	3.0
234426	cookies by design sugar shortbread cookies	20	506822	2008-04-15	 4.0	4.0	11.0	6.0
234427	cookies by design sugar shortbread cookies	20	506822	2008-04-15	 4.0	4.0	11.0	6.0
234428	cookies by design sugar shortbread cookies	20	506822	2008-04-15	 4.0	4.0	11.0	6.0

234429 rows × 23 columns

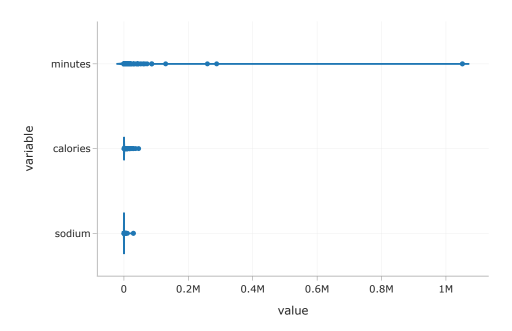
Now this code would be used later on when we need to groupby using the recipe_id column or the user_id column for different purposes. The handling for different columns are also defined as below, which is different according to waht we need the columns are for later on in the modeling process.

```
'tags':lambda x: list(chain.from_iterable(x)),
'rating':'mean',
'minutes':'mean',
'calories':'mean',
'description':lambda x: list(x),
'n_ingredients':'mean',
'ingredients':lambda x: list(chain.from_iterable(x)),
'contributor_id':lambda x: list(x),
'review':lambda x: list(x),
})
```

Univariate & Bivariate Analysis

Okay, after data cleaning, let's draw some graph to see what kind of data we are dealing with

```
In [ ]: px.violin(cleaned, x=['sodium','calories','minutes'])
```



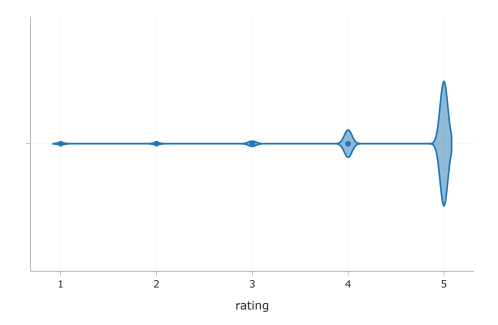
Looks like that our data have a lot of outliers! we might want to write a function to deal with that. Here we are writing the function outlier, which will be used quite often later on.

```
def outlier(df):
    '''take care of outliers in the data frame'''
    # Remove outlier in graph dierctly

check = ['minutes', 'n_steps', 'n_ingredients', 'calories', 'total_fat', 'sugar', 'sodium',
    for col in check:#df.select_dtypes(include='number'):
        q_low = df[col].quantile(0.01)
        #print(q_low)
        q_hi = df[col].quantile(0.99)
        #print(q_hi)
        df = df[(df[col] < q_hi) & (df[col] > q_low)]

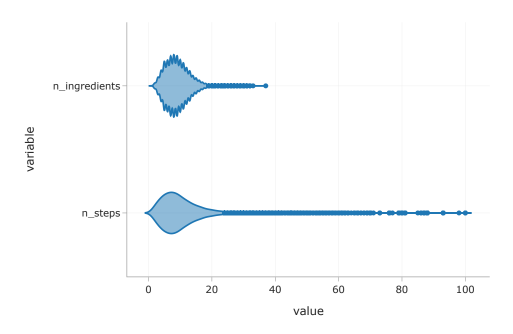
return df #same name so update df
```

```
In [ ]: px.violin(cleaned, x='rating')
```



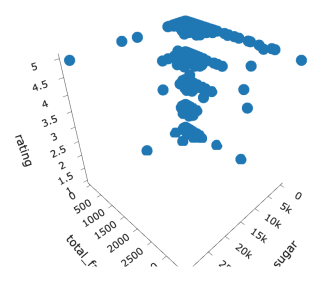
Looks like the data are kind of **unbalanced** in rating (at this point, we thought that this wouldn't effect our modle too much, but it turns out later to be one of the main challenge that we need to deal with during the moeling phase)





Seems like there is a **threshold point** for $n_{ingredients}$ and n_{steps} , this will be utilized later in our **feature engineering** section

```
In [ ]: px.scatter_3d(cleaned, x='sugar', y='total_fat', z='rating',)
```



It also seems like more sugar and more total_fat (transformed from nutrition) seems to be related to higher rating! This is quite suprising!

We actually made more edas and feature engineering with **textual features**, but we will introduce those later in the section as it is much more relevant to our modeling process.

Step 3: Assessment of Missingness

There are data missing! Why is that happening?

We are specifically working with the version of the data set that have been grouped by with recipe_id to check the missingness, each recipe_id in this case would be unique. We can start with checking which column is missing. For the easiness of graphing, we will first slice out the outliers in each of the numerical columns using outlier function, which slices out ouliers that's out of the 99th percentile of the dataset

```
In [ ]: display_df(pd.DataFrame(df.isna().sum()), 23)
```

	0
minutes	0
n_steps	0
n_ingredients	0
avg_rating	1679
rating	1679
calories	0
total_fat	0
sugar	0
sodium	0
protein	0
sat_fat	0
carbs	0
steps	0
name	0
description	48
ingredients	0
user_id	0
contributor_id	0
review_date	0
review	0
recipe_date	0
tags	0

NMAR Analysis

From the first step analysis by just observing the data and looking at the website, it seems like the column name is **Missing Completely At Random (MCAR)** as it doesn't seem to be fitting to any of the all three other catagories.

However, on the other hand, the rating column seems to be **Not Missing At Random (NMAR)** because from what the website is showing, some people just didn't give rating, so the rating itself doesn't exist during the data collection process, so it makes sense for it to be null. We manually added <code>np.NaN</code> into the data set where previously it was filled a zero in the data set. Since <code>avg_ratng</code> is calculated from using the <code>rating</code> column, <code>avg_rating</code> would then be **Missing At Random (MAR)** dependent on <code>rating</code>.

One interesting one to analyze is <code>description</code>, because it is hard to say directly how it may be correlated to any other columns in this data set, we suspect it to be <code>MAR</code>, but we will prove it to be <code>MAR</code> in the next section.

MAR Analysis

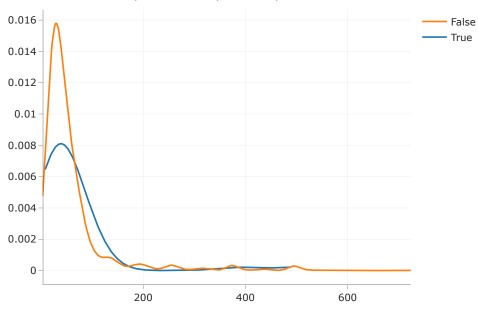
Decision Rule for description

Let's assume that the missingess of description column is related to the col column for continuous columns, wouldn't depend on discrete columns.

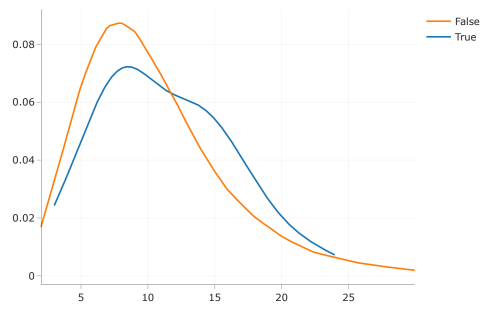
The below functions are used for conducting graphing for checking potential MAR columns and also for conducting permutation testing

```
In [ ]: def create_kde_plotly(df, group_col, group1, group2, vals_col, title=''):
            '''Create the kde plot for checking column potential dependencies'''
            fig = ff.create_distplot(
                hist_data=[df.loc[df[group_col] == group1, vals_col], df.loc[df[group_col] == group2, va
                group_labels=[group1, group2],
                show_rug=False, show_hist=False
            return fig.update_layout(title=title)
        def mar check continuous(df,miss col, dep col):
            '''Full checking mar by simulating mar data then graphing it,
            miss_col must be catagorical and dep_col must be continuous'''
            missing = df[miss_col].isna()
            df_missing = df.assign(mar_missing = missing)[['mar_missing', dep_col]]
            fig = create_kde_plotly(df_missing, 'mar_missing', True, False, dep_col, title=f'MAR Graph o
            return fig.show()
In [ ]: def permutation_ks(df, miss_col, dep_col, rep):
             '''conduct permutation testing for testing mar in data frame '''
            def permutation_test(df, rep, dep_col):
                 '''test statistics is the KS statistics'''
                # line of missing of description that may base on dep_col?
                observe = ks_2samp(df_missing.query('mar_missing')[dep_col],
                                   df_missing.query('not mar_missing')[dep_col]).statistic
                # making a distrbution where missing of description does not depend on dep_col
                n_repetitions = rep
                null = []
                for _ in range(n_repetitions):
                    with_shuffled = df.assign(shuffle = np.random.permutation(df['mar_missing']))
                    difference = ks_2samp(with_shuffled.query('shuffle')[dep_col],
                                           with_shuffled.query('not shuffle')[dep_col]).statistic
                    null.append(difference)
                return observe, null
            missing = df[miss_col].isna()
            df_missing = df.assign(mar_missing = missing)[['mar_missing', dep_col]]
            observe, null = permutation_test(df_missing, rep, dep_col)
            fig = px.histogram(pd.DataFrame(null), x=0, histnorm='probability', title=f'KS Distribution
            fig.add_vline(x=observe, line_color='red', line_width=1, opacity=1)
            p = (observe <= null).mean()</pre>
            print(f'p_value is {p}')
            return fig.show()
In [ ]: | for col in df.drop(columns=['avg_rating','rating']).select_dtypes(include='number'):
            mar_check_continuous(df, 'description', col)
```

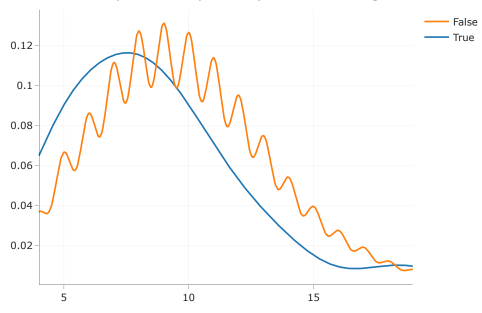
MAR Graph of description Dependent on minutes



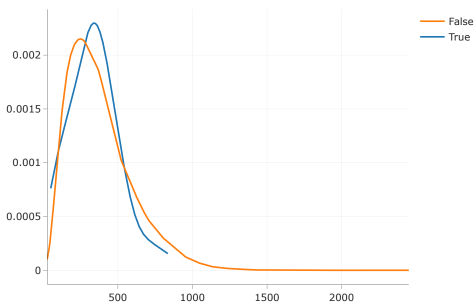
MAR Graph of description Dependent on n_steps



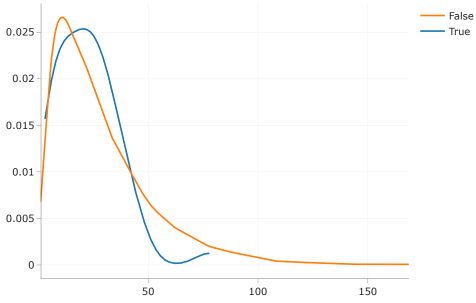
MAR Graph of description Dependent on n_ingredients



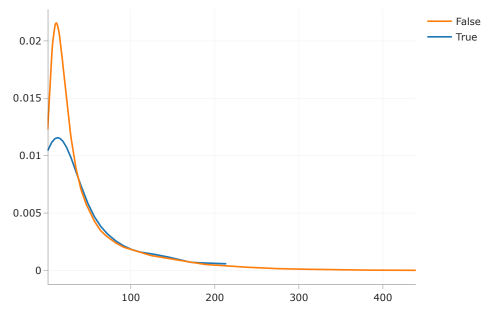
MAR Graph of description Dependent on calories

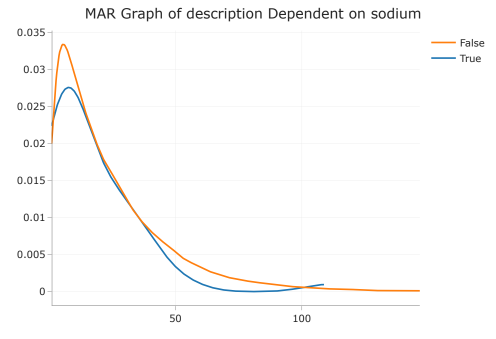


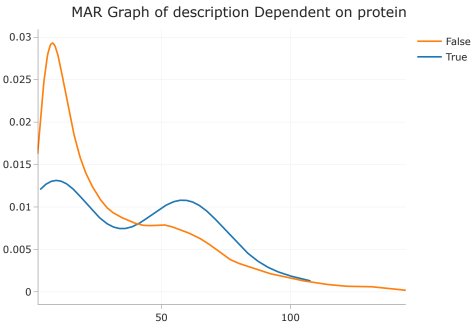


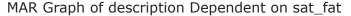


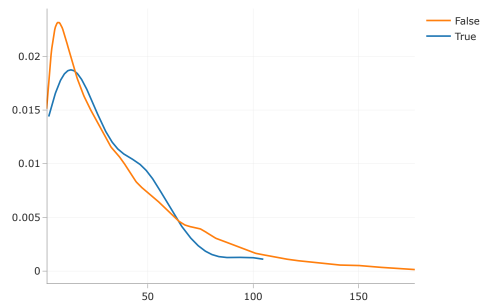
MAR Graph of description Dependent on sugar



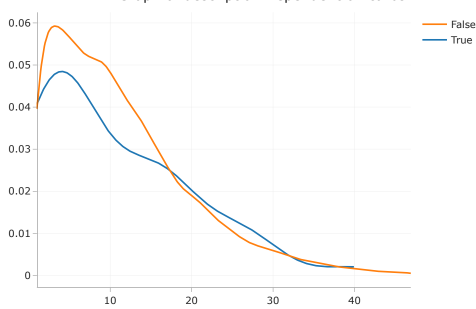








MAR Graph of description Dependent on carbs



description seems to also depend on n_ingredients. This is a very interesting graph because looks like the graph **shape** is quite different with the **mean** the same, instead of using permutation test statistics that involves **mean** we use **K-S statistics** instead (we have also down a test using differences in mean as well, which fail to identify any results).

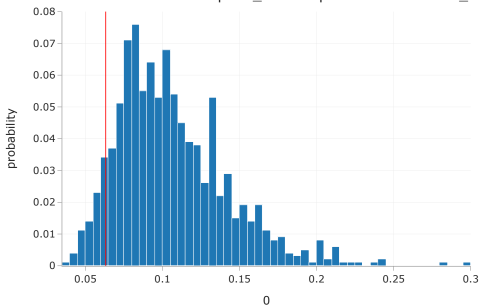
Permutation Testing Using K-S Statistics

Now we want to perform permutation testing with each of the continuous variable within the data set (assuming that the missingness of description depends on them) and plot the distribution

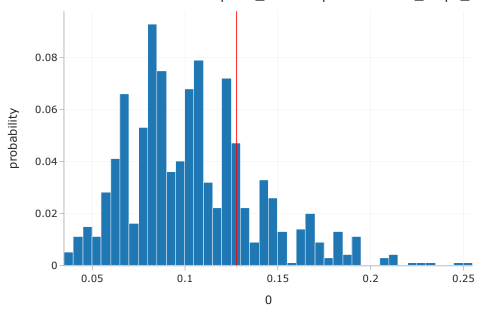
We decide to use a testing threshold of $p=0.05\,$

```
In [ ]: for col in df.drop(columns=['avg_rating','rating']).select_dtypes(include='number'):
    permutation_ks(df, 'description', col, 1000)
```

KS Distribution for Null description_col is dependent on minutes_col

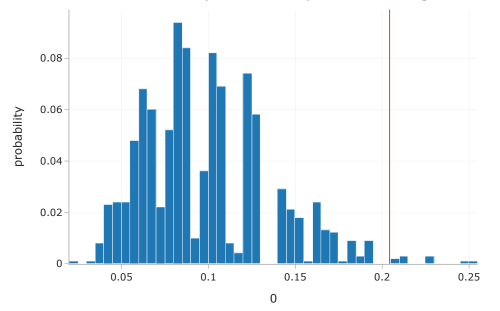


 $\rm p_value\ is\ 0.217$ KS Distribution for Null description_col is dependent on n_steps_col

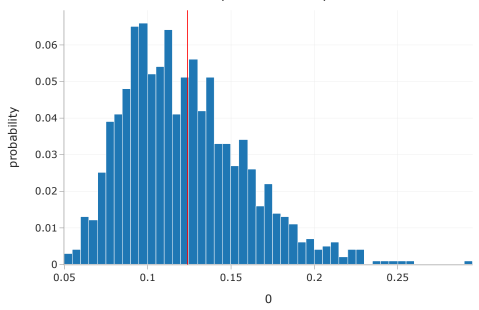


p_value is 0.01

5 Distribution for Null description_col is dependent on n_ingredients_c

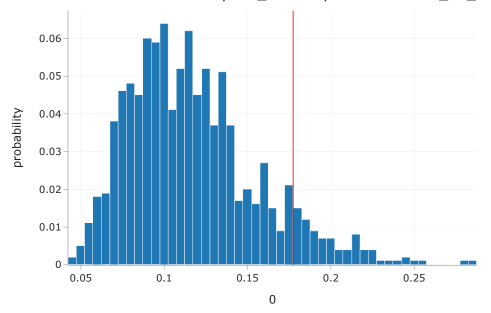


p_value is 0.431 KS Distribution for Null description_col is dependent on calories_col

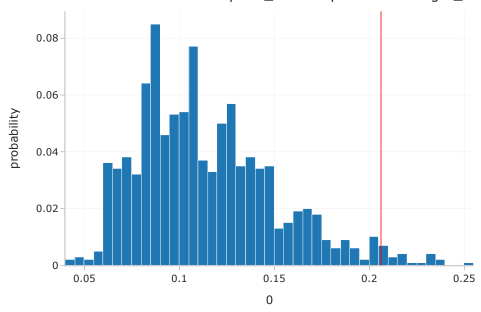


p_value is 0.083

KS Distribution for Null description_col is dependent on total_fat_col

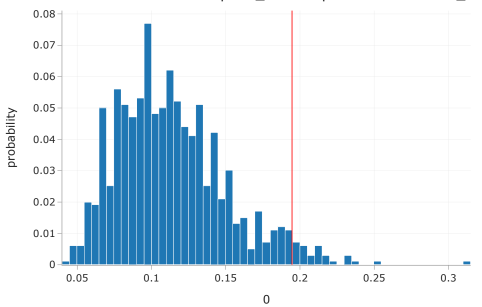


p_value is 0.022
KS Distribution for Null description_col is dependent on sugar_col

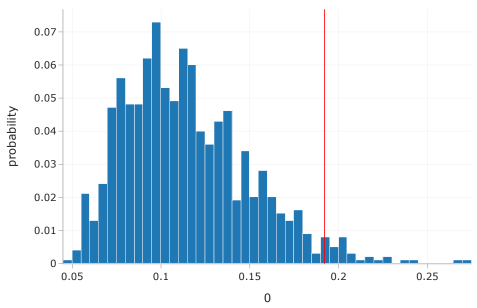


p_value is 0.037

KS Distribution for Null description_col is dependent on sodium_col

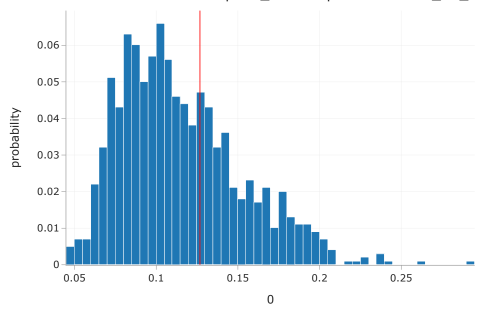


 $\rm p_value\ is\ 0.032$ KS Distribution for Null description_col is dependent on protein_col

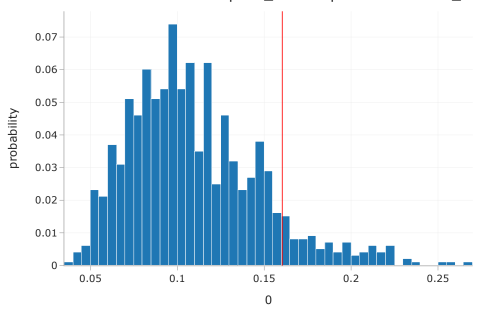


p_value is 0.343

KS Distribution for Null description_col is dependent on sat_fat_col



p_value is 0.092
KS Distribution for Null description_col is dependent on carbs_col



- seems like missingess for description is related to n_ingredients
- seems like missingness in description is not related to calories or n_steps

Step 4: Hypothesis Testing

For this section, we will be working with the same data frame that was used in the missingness mechanism section, so a data frame that is grouped by recipe_id.

Since we want to do certain textual feature analysis for our predictive model, we were wondering whether TF
IDF of the description columns would actually play a role in deternmining the rating of an recipe. This
can be deemed as a mini-warmup for our modeling procedure later on.

Term Frequency Inverse Document Frequency

TF-IDF is a very naive but common and well performing technique that people use to understand textual features. It essentially meausres the **how important** an word t is for an sentence in comparison with all sentences in the document. The TF-IDF Formula is a as follows:

$$\begin{split} \text{tfidf}(t,d) &= \text{tf}(t,d) \cdot \text{idf}(t) \\ &= \frac{\text{$/$\# of occurrences of t in d}}{\text{total $/$\# of words in d}} \cdot \log \bigg(\frac{\text{total $/$\# of documents}}{\text{$/$\# of documents in which t appears}} \bigg) \end{split}$$

We will be using TfidfVectorizer to help our calculation

```
In [ ]: # import here first as it is usef for computing TF-IDF
from sklearn.feature_extraction.text import TfidfVectorizer
```

In here we are just splitting the data frame into high_score and low_score

```
In []: df = df[['name','description','tags','steps','ingredients','contributor_id','rating']] # avg_rat
    df_high = df[df['rating']>=4]
    df_low = df[df['rating']<4]

lst_high = df_high['description'].explode().astype(str)
lst_low = df_low['description'].explode().astype(str)</pre>
```

In this step we are first using the TfidfVectorizer from sk_learn to compute the TF-IDF table

Differences in Max for TF-IDF

We want to see whether the distribution of high_rated recipes and the distribution of low_rated recipes actually come from the same distribution. Thus, we will be performing a **permutation test** here with the following hypothesis:

- **Null hypothesis**: There **are no** differences in the distribution for the **high_rated** recipes and low_rated` recipes.
- Alternative hypothesis: There are differences in the distribution for the high_rated recipes and low_rated`recipes.

We decide to use a testing threshold of $p=0.05\,$

As for the **test statistics**, we actually have many options, but they all circles around the **differences** of something:

- Using sum -> longer sentences have greater sum
- Using mean -> very easy to be influenced by outlier
- Using partial-mean -> get the most essence part of the sentence, however, complexity too high because of the sorting
- Using max -> most important one word's TF-IDF

This section provide a solid prove of why we are using TF-IDF as a feature for our predictive model!

```
In []: tfidf_max_high = high_tfidf.max(axis=1)
    tfidf_max_low = low_tfidf.max(axis=1)

max_high = df_high.reset_index().assign(tfidf = tfidf_max_high, good=True)
    max_low = df_low.reset_index().assign(tfidf = tfidf_max_low, good=False)

big_df = pd.concat([max_high, max_low], axis=0)
    big_df
```

Out[]:		recipe_id	name	description	tags	 contributor_id	rating	tfidf	good
	0	275030.0	paula deen s caramel apple cheesecake	thank you paula deen! hubby just happened to	[60- minutes-or- less, time- to-make, course, pre	 [666723, 666723, 666723, 666723, 666723, 66672	5.0	0.28	True
	1	275033.0	penne with bacon spinach mushrooms	from woman's day magazine.	[bacon, 30- minutes-or- less, time- to-make, cour	 [166642]	5.0	0.72	True
	2	275036.0	easy weeknight corn	i threw some things together in a dutch oven a	[15-minutes- or-less, time-to- make, course, mai	 [590640, 590640]	5.0	0.33	True
	3642	535783.0	cheesesteak stuffed onion rings	surprise your family and friends with an onion	[60- minutes-or- less, time- to-make, main- ingred	 [33186, 33186]	3.0	0.36	False
	3643	536688.0	coco oatmeal honey cookies	the fiber-rich cookies are good for snacking.h	[30- minutes-or- less, time- to-make, course, pre	 [2002170767, 2002170767]	3.0	0.37	False
	3644	536843.0	sheet pan turkey caprese meatballs with rosema	description: try these turkey caprese meatball	[60- minutes-or- less, time- to-make, course, mai	 [2001112113, 2001112113]	3.0	0.36	False

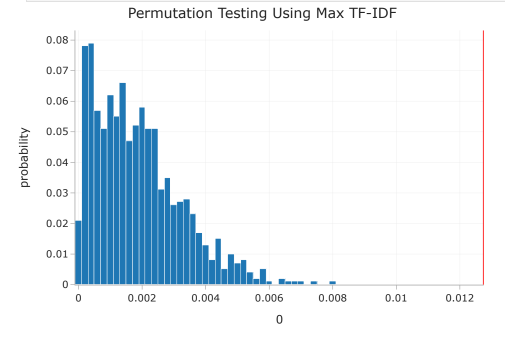
54873 rows × 10 columns

Permutation Testing

```
In []: observe = big_df.groupby('good')['tfidf'].mean().diff().abs().iloc[-1]

# making a distrbution where missing of description does not depend on dep_col
n_repetitions = 1000
null = []
for _ in range(n_repetitions):
    with_shuffled = big_df.assign(shuffle = np.random.permutation(big_df['good']))
    difference = with_shuffled.groupby('shuffle')['tfidf'].mean().diff().abs().iloc[-1]
    null.append(difference)
```

fig = px.histogram(pd.DataFrame(null), x=0, histnorm='probability', title=f'Permutation Testing
fig.add_vline(x=observe, line_color='red', line_width=1, opacity=1)



The result is significant! We reject the null hypothesis! There is a difference in the distribution for high rated recipes and low rated recipes.

Step 5: Framing a Prediction Problem

From the previous section we have learned that Recipe's Max TF-IDF distribution is different for high_rated recipe than low_rated recipe, so now we want to go a step further: we want to predict rating as a classfication problem to demonsrate user preference and as a potential prior to recommander system

Specifically, we want to predict rating (5 catagories) in the original data frame to demonstarte understanding of user preference. In this section we will be using the original big DataFrame for predicting rating.

Step 6: Baseline Model

Just to keep everything clear, we recalled all the cleaning function here and some neccessary extraction performed

```
In []: # for modeling transformation
    from sklearn.preprocessing import FunctionTransformer, OneHotEncoder, Binarizer, RobustScaler
    from sklearn.base import BaseEstimator, TransformerMixin
    from sklearn.compose import Pipeline
    from sklearn.compose import ColumnTransformer
    from sklearn.feature_extraction.text import CountVectorizer
    from sklearn.feature_extraction.text import TfidfVectorizer
    from sklearn.decomposition import PCA

# for modeling hyperparameter tuning
    from sklearn.model_selection import train_test_split
    from sklearn.model_selection import GridSearchCV, RandomizedSearchCV
    from sklearn.ensemble import RandomForestClassifier
    from sklearn.dummy import DummyClassifier
```

Out[]:		n_ingredients	minutes	n_steps	description		recipe_date	review	is_good	is_low
	0	9	40	10	these are the most; chocolatey, moist, rich, d	•••	2008-10-27	These were pretty good, but took forever to ba	False	False
	1	11	45	12	this is the recipe that we use at my school ca		2011-04-11	Originally I was gonna cut the recipe in half	True	False
	2	9	40	6	since there are already 411 recipes for brocco		2008-05-30	This was one of the best broccoli casseroles t	True	False
	•••									
	234426	7	20	5	i've heard of the 'cookies by design' company,		2008-04-15	This recipe tastes nothing like the Cookies by	False	True
	234427	7	20	5	i've heard of the 'cookies by design' company,	•••	2008-04-15	yummy cookies, i love this recipe me and my sm	True	False
	234428	7	20	5	i've heard of the 'cookies by design' company,		2008-04-15	I work at a Cookies By Design and can say this	False	False

154763 rows × 15 columns

Handling Missing Data

It have been shwon earlier that the missingness of the rating columns seems to be **NMAR**, so it is not dependent on the column but rather depending on itself. Thus, we will be imputing the ratings through **random imputation**.

• This may cause more rating of 5 to come up

```
In []: def prob_impute(s):
    s = s.copy()
    num_null = s.isna().sum()
    fill_values = np.random.choice(s.dropna(), num_null)
    s[s.isna()] = fill_values
```

```
return s
base_df['rating'] = prob_impute(base_df['rating'])
base_df = base_df.dropna()
```

Train/Validate/Test Split

We are splitting the main data set into 3 components of train, validate, and test. The main data set is plit to big_train and test first with big_train being 75% of the data. Then, the big_train data set is split again into the validate and the actual train data set with 75% in the train data set again. Each set is taking the percentage as calculated below:

```
Test: 25%Train_big: 75%Validate: 18.75%Train: 56.25%
```

```
In []: X = base_df.drop('rating', axis=1)
y = base_df['rating']
X_big_train, X_test, y_big_train, y_test = train_test_split(X, y, test_size=0.25, random_state=1
X_train, X_val, y_train, y_val = train_test_split(X_big_train, y_big_train, test_size=0.25, random_state=1)
```

Feature Engineering & Transformations

```
In [ ]: class StdScalerByGroup(BaseEstimator, TransformerMixin):
            '''takes in two separate, fitting data may not be transforming data (training)'''
            def __init__(self):
                pass
            def fit(self, X, y=None):
                '''fit using one type of data'''
                # X might not be a pandas DataFrame (e.g. a np.array)
                df = pd.DataFrame(X)
                # Compute and store the means/standard-deviations for each column (e.g. 'c1' and 'c2'),
                mean_group = df.groupby(df.columns[0]).mean()
                std_group = df.groupby(df.columns[0]).std()
                for col in mean_group:
                    mean_group = mean_group.rename(columns={col:f'{col}_mean'})
                    std_group = std_group.rename(columns={col:f'{col}_std'})
                self.grps_ = pd.concat([mean_group,std_group],axis=1)
                return self
            def transform(self, X, y=None):
                '''may be different data'''
                    getattr(self, "grps_")
                except AttributeError:
                    raise RuntimeError("You must fit the transformer before transforming the data!")
                def standardize(x, col):
                    group = x.name
                    mean = self.grps_.loc[group, f'{col}_mean']
                    std = self.grps_.loc[group, f'{col}_std']
                    norm = (x - mean) / std
                    return norm
                df = pd.DataFrame(X)
                new=pd.DataFrame()
                for col in df.columns[1:]:
```

Baseline Model's Pipeline

In the basic model pipeline we are working with not a great number of features:

- 1. binarized n_step with threshold 25
- 2. binarized n ingredients with threshold 20
- 3. normalized minutes with respects to binarized n_steps using the customized class StdScalerByGroup
- 4. normalized minutes with respects to binarized n_ingredients using the customized class as above
- 5. simple counts of tags column

The pipeline for the model is constituted with a simple **Random Forest** multi-class classfier with hyperparameter tuned

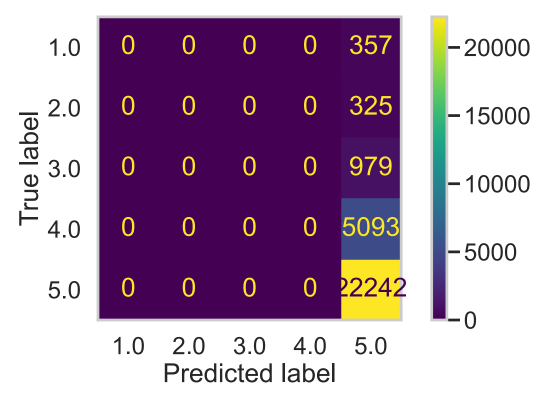
```
In [ ]: norm_relative = Pipeline([
            ('bi_nsteps', Binarizer(threshold=25)),
            ('norm_minutes_binary_nsteps', FunctionTransformer(lambda x: StdScalerByGroup().fit(x).trans
        ])
        preproc_rf = ColumnTransformer(
            transformers=[
                ('bi_nsteps', Binarizer(threshold=25),['n_steps']),
                ('bi_ningredients', Binarizer(threshold=20),['n_ingredients']),
                ('norm_minutes_binary_nsteps',norm_relative,['n_steps','minutes']),
                ('norm_minutes_binary_ningredients',norm_relative,['n_ingredients','minutes']),
                ('tag_counts', FunctionTransformer(tag_counts), ['tags']),
            1,
            remainder='drop'
        pl_base = Pipeline([
            ('preprocessor', preproc_rf),
            ('rfc', RandomForestClassifier(max_depth=2,
                                            n_estimators=140,
                                            criterion='entropy'
                                            min_samples_split=2))
        ])
```

```
In [ ]: pl_base.fit(X_train, y_train)
pl_base.score(X_val, y_val)
```

Out[]: 0.7670713201820941

This looks pretty good! Roughly 77% on validation set is pretty good! But let's dig deeper first

```
In [ ]: ConfusionMatrixDisplay.from_estimator(pl_base, X_val, y_val)
   plt.grid(False)
```



```
In []: base_df['rating'].value_counts() / base_df.shape[0]
Out[]: 5.0     0.77
     4.0     0.18
     3.0     0.03
     1.0     0.01
     2.0     0.01
```

Turns out the original dataset is highly **unbalanced**, making the model always predicting a rating of 5 not missing many of the other details. This also means that as long as the model is always predicting the rating of 5, it will get an accuracy of 77% because 77% of the rating is 5 -> accuracy doesn't entell everything!. Thus, we need a better model than this that can capture some what more feature information, more engineering is needed!

Step 7: Final Model

Name: rating, dtype: float64

"A good model is the combination of model section" + feature engineering + hyperparameter tuning"*

Now with the previous baseline model's problem in mind, let's make some actual useful feature engineering, mainly we will be utilizing these features:

The previous features are carried over to this model, which includes:

- 1. binarized n_step with threshold 25, this is a result from eda
- 2. binarized n_ingredients with threshold 20, this is a result from eda
- 3. normalized minutes with respects to binarized n_steps
- 4. normalized minutes with respects to binarized n_ingredients
- 5. simple counts of tags column, showing how many tags are in each tag column

In addition, awe also added afew more features to capture the relationship we saw from EDA, which includes:

- Some numerical columns of sugar, sodium, calories, total_fat that have being standerlized using RobustScaler
- 2. Two TF-IDF that have been one hot encoded:
 - In particular, the naive approach is to use the highest TF-IDF for each of the words are extracted for each of the sentence using argmax, representing the most important words in a sentence (we are using argmax here is for considering the complexity of this model, later implementations can utilize more words that have high TF-IDF)
 - We then construct a pool of highest TF-IDF words in the **low** rating dataset, which was originally defined as rating lower than or equal to 3 and it is stored as a boolean indicator in the is_low column.
 - Finally, we want to see whether or not the current sentence's highest TF-IDF word is in such pool of words
 - We perform such operations with both the name column and also the description column
 - Remark: this feature improved the final model by roughly 10% accuracy, this is the detect_key_low(df) function
 - We ahve also tried to trade off some complexity with better accuracy by using the count of the 5 top TF-IDF words in each row (just this function runs for about 3m)
 - However, the performance didn't perform as well as argmax, whihch may be due to extra noise added (48% accuracy with 5 words and 50% accuracy with one word)
- 3. The recipe_dtae column have also being taken out with only the year of the recipe and then one hot encoded as well.
- 4. At last, we also used the tag column of each of the sentence to perform one hot encoding
 - We first performed one hot encoding to transform each tag to a numerical boolean representation.
 However, this makes the feature space to reahc to about 500 features, which adds too much sparsity to the feature space and may introduces noises
 - Thus we filtered out all the irrelevant or low counted tags (<1000 counts) and reduces teh feature spac to only adding 80 more features
 - At last, we conducted pca to reduce the adding feature space to just abou 10 features and this value seems to work well with the data set experimentally.
 - The tag_ohe_pca(df) function takes care of this step
- 5. Analyzing whether the review columns contain certain sentiment words in it, evaluated by the is_sentiment(df) function
- 6. We have taken out irrelevant features such as the naive_bayes encoder taht we have implemented

```
In [ ]: def tag counts(df):
            '''number of tags counted'''
            return pd.DataFrame(df['tags'].apply(lambda x: len(x)).rename('counts'))
        def detect key low(df):
            '''transforming description's tfidf to actual most important word in a description then comp
            def key largest(row):
                return row.index[row.argmax()] #[row.argsort()][-5:]
            def make tfidf(series):
                lst = series.explode().astype(str).values # this may be slow
                count = TfidfVectorizer()
                count.fit(lst)
                return pd.DataFrame(count.transform(lst).toarray(), columns=count.get_feature_names_out(
            tfidf_low = make_tfidf(df[df['is_low']==True][df.columns[1]])
            tfidf_base = make_tfidf(df[df.columns[1]])
            keyword_all = tfidf_base.apply(key_largest, axis=1) #argmax a bit faster
            keyword_low = tfidf_low.apply(key_largest, axis=1)
            pool_low = keyword_low.unique() #.explode().unique()
            in_low = keyword_all.apply(lambda x: x in pool_low) #.apply(lambda x: sum([word in pool_low)
```

```
return pd.DataFrame(in_low)
def tag_ohe_pca(df):
    '''OHE all the tag result after it have being pca dimension reduced to 50'''
   # getting all the unique one quick
   set = [j for i in df['tags'].tolist() for j in i] # explode in a time complexity efficient w
   count = CountVectorizer()
   count.fit(set).transform(set)
   my dict = np.array(list(count.vocabulary .keys()))
   def helper function(list, dict):
        return np.array([i in list for i in dict])
   a = df["tags"].apply(lambda x:helper_function(x, my_dict))
   # change array of array into 2D array
   df_pca = pd.DataFrame(data = np.stack(a.to_numpy()),columns=my_dict)
   flipped = df_pca.T
   filter_df = flipped[flipped.sum(axis=1)>1000].T # keep only useful tags
   # conduct PCA to reduce to just 50 dimensions
   pca = PCA(n_components=10)
   reduced = pca.fit_transform(filter_df)
   return reduced
def is_sentiment(df):
    '''For detecting sentiment words in the review column'''
   word_list = ['awful', 'fav', 'well',
             'yet', 'fantastic',
             'pretty good', 'dislike', 'hate', 'bad',
             'delicious', 'wonderful',
             'great', 'but', 'good', 'next',
             'excellent', 'nice', 'bland', 'maybe',
             'loved', 'sorry', 'think', 'however', 'would',
             'perfect', 'very', 'keeper', 'liked', 'made']
   out = df['review'].apply(lambda x: word in x for word in word list)#.sum(axis=1)
   return pd.DataFrame(out.astype(int))
```

Final Model's Pipeline

Since this is a **multi-class classifictaion** problem and the data is also highly **unbalanced**, we are also adding a **dummy** classifier that classifies uniformally at random to bench mark our modle's performances. Of course, we will also use different evaluation metrics later to demonstarte the model's performances as well, the dummy classifier is just an "easy to view" example.

```
In []: # dummy classfier with uniformally selections
dummy_clf = DummyClassifier(strategy="uniform")
dummy = dummy_clf.fit(X_train, y_train)
```

For the pipeline, we are still doing an **Homogenous Ensemble Learning** with decision tree as it have being shown to perform the best experimentally (we have tried hetergenous ensemble learning using voting/stacking with models such as SVM and logistic regression, but none of the perform as well as the simple random forest).

We balanced the dataset by using automatic balaning argumnet "balanced", we have also tried to use self customized dictionaries for assigning weights, However, this wouldn't be generalizable to unseen data as the

distribution of data changes. The sk_learn packages does automatic weight assigning by the following formula:

```
n_samples/(n_classes*np.bincount(y))
```

This model pipeline takes about 30 seconds to fit

```
In [ ]: norm_relative = Pipeline([
            ('bi_nsteps', Binarizer(threshold=25)),
            ('norm_minutes_binary_nsteps', FunctionTransformer(lambda x: StdScalerByGroup().fit(x).trans
        1)
        key ohe = Pipeline([
            ('tfidf',FunctionTransformer(detect_key_low)),
            ('key_ohe', OneHotEncoder(drop='first'))
        1)
        date_ohe = Pipeline([
            ('date_get', FunctionTransformer(lambda x: pd.DataFrame(x['recipe_date'].dt.year))),
            ('date_ohe', OneHotEncoder())
        ])
        preproc_rf = ColumnTransformer(
            transformers=[
                ('tfidf_key_ohe_description', key_ohe, ['is_low', 'description']),
                ('tfidf_key_ohe_name', key_ohe, ['is_low', 'name']),
                ('bi_nsteps', Binarizer(threshold=25),['n_steps']),
                ('bi_ningredients', Binarizer(threshold=20),['n_ingredients']),
                ('norm_minutes_binary_nsteps',norm_relative,['n_steps','minutes']),
                ('norm_minutes_binary_ningredients',norm_relative,['n_ingredients','minutes']),
                ('tag_counts',FunctionTransformer(tag_counts),['tags']), # good for predict 5
                ('robust_nutrition',RobustScaler(),['sugar','sodium','calories','total_fat']),
                ('date_ohe', date_ohe,['recipe_date']), # can't be too sparse, 10 max
                ('tag_pca',FunctionTransformer(tag_ohe_pca),['tags']),
                ('is_sentiment',FunctionTransformer(is_sentiment),['review'])
            ],
            remainder='drop'
        class_weight= 'balanced' # balanced weighting automatically calculates th best weighting!
        pl rf = Pipeline([
            ('preprocessor', preproc rf),
            ('rfc', RandomForestClassifier(max depth=18,
                                           n estimators=130,
                                            criterion='entropy',
                                            min samples split=2,
                                            class weight=class weight
                                            ))
        ])
        pl_rf.fit(X_train, y_train)
```

```
Out[]: Pipeline(steps=[('preprocessor',
                          ColumnTransformer(transformers=[('tfidf_key_ohe_description',
                                                             Pipeline(steps=[('tfidf',
                                                                               FunctionTransformer(func=<fu
         nction detect_key_low at 0x15ef414c0>)),
                                                                              ('key_ohe',
                                                                               OneHotEncoder(drop='firs
        t'))]),
                                                             ['is_low', 'description']),
                                                            ('tfidf_key_ohe_name',
                                                             Pipeline(steps=[('tfidf',
                                                                               FunctionTransformer(func=<fu
        nction detect_key_low at 0x...
                                                                               FunctionTransformer(func=<fu</pre>
        nction <lambda> at 0x15fa831f0>)),
                                                                              ('date_ohe',
                                                                               OneHotEncoder())]),
                                                             ['recipe_date']),
                                                            ('tag pca',
                                                             FunctionTransformer(func=<function tag_ohe_pc</pre>
        a at 0x15ef410d0>),
                                                             ['tags']),
                                                            ('is_sentiment',
                                                             FunctionTransformer(func=<function is sentime</pre>
        nt at 0x15ef41670>),
                                                             ['review'])])),
                          ('rfc',
                          RandomForestClassifier(class weight='balanced',
                                                   criterion='entropy', max_depth=18,
                                                   n estimators=130))])
```

Hyperparameter Tunning

We have performed Grid Search and Random Search for the best parameters for teh Random Forest Classifier. However, for the complexity of running this notebook, we only tuned the model once and then turned this cell off.

Model Evaluation

To really understand what we are evaluating, we need to first understand what metrics matters to us:

Example:

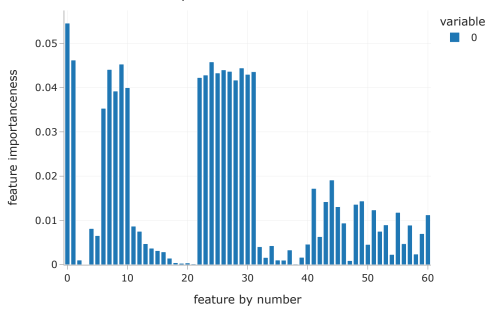
- Precision for 5: Out of all 5 we predicted, how many are actually 5
- Recall for 5: Out of all actual 5, how many did we get right

We care about getting a correct rating for recommandation, we care about finding **Recall** but still considering precision, accuracy, F1 scores

Feature Importantness Analysis

```
In []: feature = pd.DataFrame(pl_rf.named_steps['rfc'].feature_importances_)
    fig = px.bar(feature, title='Feature Importantness for Random Forest', labels={'value':'feature fig.show()
```

Feature Importantness for Random Forest

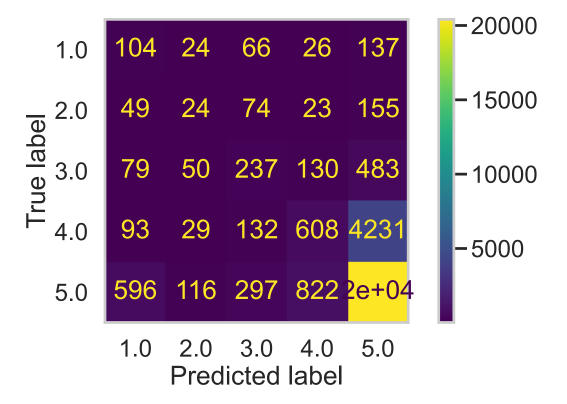


We have 60 features in our model with feature 0 and feature 1 having the most effect! these are the 2 argmax

TF-IDF encoder that we have implemented, this is consistent with ou previous **permutation testing** that shows the TF-IDF distribution for high_rated and low_rated recipes are different distributions

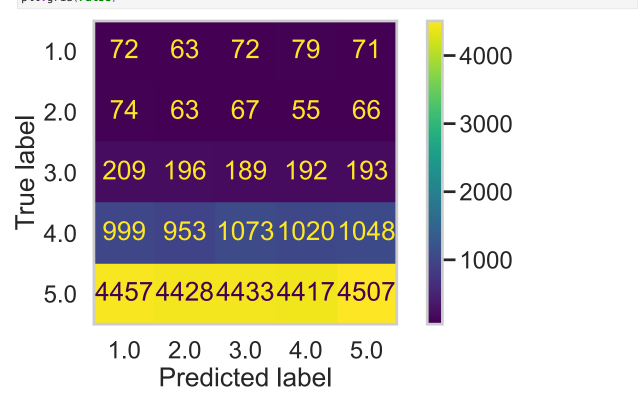
Confusion matrix for random forest classifier

```
In [ ]: ConfusionMatrixDisplay.from_estimator(pl_rf, X_val, y_val)
   plt.grid(False)
```



Confusion matrix for dummy classifier

In []: ConfusionMatrixDisplay.from_estimator(dummy, X_val, y_val)
 plt.grid(False)



Accuracy for random forest classifier

In []: pl_rf.score(X_val, y_val)

Out[]: 0.7378259070216582

Accuracy for dummy classifier

```
In [ ]: dummy.score(X_val, y_val)
```

Out[]: 0.19813077665884948

0u

Full metrics for random forest classfier

In []: metrics_all(pl_rf, X_val, y_val)

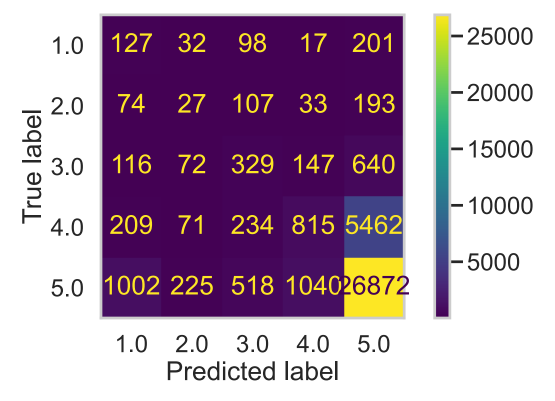
t[]:		precision	recall	f1_score	count
	1	0.29	0.11	0.16	919.0
	2	0.08	0.11	0.09	243.0
	3	0.24	0.29	0.27	806.0
	4	0.12	0.39	0.18	1587.0
	5	0.92	0.80	0.86	25441.0

Full metrics for dummy classifier

Out[]:		precision	recall	f1_score	count
	1	0.21	0.01	0.02	5840.0
	2	0.19	0.01	0.02	5752.0
	3	0.20	0.03	0.06	5825.0
	4	0.20	0.17	0.19	5864.0
	5	0.20	0.76	0.31	5715.0

Testing on Actual Test Data Set

```
In [ ]: ConfusionMatrixDisplay.from_estimator(pl_rf, X_test, y_test)
    plt.grid(False)
```



In []:	pl	_rf.score	(X_test	y_test)		
Out[]:	0.	0.7273221075502444					
in []:	me	trics_all	(pl_rf,	X_test,	y_test)		
ut[]:		precision	recall	f1_score	count		
	1	0.28	0.09	0.13	1534.0		
	2	0.07	0.07	0.07	444.0		
	3	0.25	0.26	0.26	1262.0		
	4	0.13	0.39	0.19	2216.0		
	5	0.90	0.81	0.85	33205.0		

Step 8: Fairness Analysis

We want to evaluate whether the model is fair for treating all populations. In particular, we want to check in the scope of looking at the predictions for the vegan group and the vegetarian group. Let's first check how many of them are in the data set.

```
In []: X_test['tags'].apply(lambda x: 'vegetarian' in x).sum()
Out[]: 5683
In []: X_test['tags'].apply(lambda x: 'vegan' in x).sum()
Out[]: 1432
In []: out = X_test.assign(prediction = pl_rf.predict(X_test))
In []: is_in_tag = out['tags'].apply(lambda x: ('vegetarian' in x)|('vegan' in x))
```

```
out = out.assign(is_in = is_in_tag)
```

Let's check the grouped by mean first

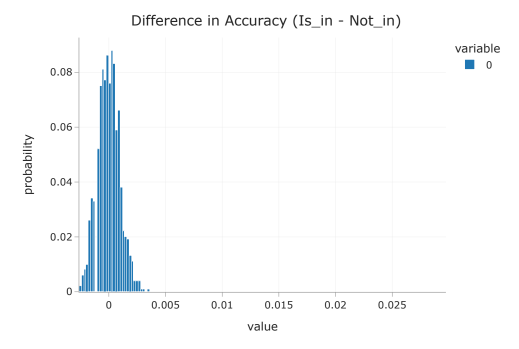
```
In []: out.groupby('is_in')['prediction'].mean()
Out[]: is_in
    False     4.67
     True     4.74
     Name: prediction, dtype: float64
```

Difference Significant?

We run a **permutation test** to see if the difference in accuracy is significant.

- **Null Hypothesis**: The classifier's accuracy is the same for both vegan + vegetarian tags and non vegan + vegetarian tags, and any differences are due to chance.
- Alternative Hypothesis: The classifier's accuracy is higher for non vegan + vegetarian tags.
- Test statistic: Difference in accuracy (is_in minus not_in).
- Significance level: 0.05

```
In [ ]: compute accuracy = lambda x: metrics.accuracy score(x['is in'], x['prediction'])
        obs = out.groupby('is_in').apply(compute_accuracy).diff().iloc[-1]
        diff_in_acc = []
        for _ in range(1000):
            s = (
                out[['is_in', 'prediction']]
                .assign(shuffle=np.random.permutation(out['is_in']))
                .groupby('shuffle')
                .apply(compute_accuracy)
                .diff()
                .iloc[-1]
            diff_in_acc.append(s)
In [ ]: fig = pd.Series(diff_in_acc).plot(kind='hist', histnorm='probability',
                                    title='Difference in Accuracy (Is_in - Not_in)')
        fig.add_vline(x=obs, line_color='red')
        fig.show()
```



```
In [ ]: (obs <= diff_in_acc).mean()</pre>
```

Out[]: 0.0

The result is **significant**, we reject the null hypothesis!