

Flatiron Phase 3 Project

Aaron Galbraith

Submitted: July 12, 2023

Business Understanding

A number of organizations install and manage water wells throughout the country of Tanzania to provide potable water to the population. Some of these wells become non-functional from time to time and need to be repaired or replaced. The government of Tanzania funds over 9,000 of these wells (about 15% of them).

Our analysis can help the government understand trends that are associated with wells that become non-functional. This can help the Tanzanian Government primarily in two ways.

1. The government can consider implementing changes in *how* (and *where*) they set up future wells so that wells have a better chance of remaining functional for a longer time without needing repair or replacement.

Of course, the government's broader goal is presumably to use their resources as efficiently as possible in delivering potable drinking water to as many residents who need it as possible. For this reason, the government surely must make choices according to some constraints that are beyond the bounds of our analysis, such as the availability or cost of certain installation options or the location choices for well placement. It could be that our recommendations for optimal well performance are incompatible with the conditions the government must deal with. For example, we may tell them to avoid installing wells in certain areas, but that might result in depriving people of water in those areas.

2. In the cases where these changes cannot be implemented, the government can more closely monitor wells that are likely to be at risk for needing repair or replacement.

It should be noted that we will eventually face a decision as to which type of error we consider the worse one in this exploration, false positives or false negatives. While neither type of error is desirable, we believe that false negatives — misidentifying non-functional wells as functional — is the worse type, as it results in depriving people of drinking water. False positives — misidentifying functional wells as non-functional — results in unneeded attention and resources being devoted to wells that are already serving effectively.

Data Understanding

Import files

Here we'll import all the tools we'll need.

```
In [1]: import numpy as np
import pandas as pd
import matplotlib.pyplot as plt
%matplotlib inline
plt.style.use('ggplot')

from itertools import combinations

import seaborn as sns

from sklearn import tree
from sklearn.model_selection import train_test_split, GridSearchCV, cross_val_score
from sklearn.linear_model import LogisticRegression
from sklearn.tree import DecisionTreeClassifier
from sklearn.metrics import accuracy_score, roc_curve, auc, confusion_matrix, classification_report
from sklearn.preprocessing import OneHotEncoder
from sklearn.ensemble import BaggingClassifier, RandomForestClassifier, AdaBoostClassifier

from scipy import stats
from scipy.interpolate import make_interp_spline

from xgboost import XGBClassifier

SEED = 10
```

The data package includes four documents. We'll open these and briefly examine their contents one at a time.

```
In [2]: df = pd.read_csv('../data/training_set_values.csv')
# show row and column counts
df.shape
```

```
Out[2]: (59400, 40)
```

```
In [3]: # show first few records
df.head()
```

```
Out[3]:
```

	id	amount_tsh	date_recorded	funder	gps_height	installer	longitude	latitude	wpt_name	num_private
0	69572	6000.0	2011-03-14	Roman	1390	Roman	34.938093	-9.856322	none	0
1	8776	0.0	2013-03-06	Grumeti	1399	GRUMETI	34.698766	-2.147466	Zahanati	0
2	34310	25.0	2013-02-25	Lottery Club	686	World vision	37.460664	-3.821329	Kwa Mahundi	0
3	67743	0.0	2013-01-28	Unicef	263	UNICEF	38.486161	-11.155298	Zahanati Ya Nanyumbu	0
4	19728	0.0	2011-07-13	Action In A	0	Artisan	31.130847	-1.825359	Shulenii	0

5 rows × 40 columns

The file "training_set_values" has 59,400 records (the wells) with 40 columns of details about those wells.

```
In [4]: df = pd.read_csv('../data/training_set_labels.csv')
# show row and column counts
df.shape
```

```
Out[4]: (59400, 2)
```

```
In [5]: # show first few records  
df.head()
```

Out[5]:

	id	status_group
0	69572	functional
1	8776	functional
2	34310	functional
3	67743	non functional
4	19728	functional

```
In [6]: # show value counts for the target variable  
df.status_group.value_counts()
```

```
Out[6]: functional           32259  
non functional        22824  
functional needs repair    4317  
Name: status_group, dtype: int64
```

The file "training_set_labels" just tells what the status of those 59,400 wells is, i.e. whether the well is functional.

```
In [7]: df = pd.read_csv('../data/test_set_values.csv')  
# show row and column counts  
df.shape
```

```
Out[7]: (14850, 40)
```

```
In [8]: # show first few records  
df.head()
```

Out[8]:

	id	amount_tsh	date_recorded	funder	gps_height	installer	longitude	latitude	wpt_name	num_private
0	50785	0.0	2013-02-04	Dmdd	1996	DMDD	35.290799	-4.059696	Dinamu Secondary School	
1	51630	0.0	2013-02-04	Government Of Tanzania	1569	DWE	36.656709	-3.309214	Kimnyak	
2	17168	0.0	2013-02-01	NaN	1567	NaN	34.767863	-5.004344	Puma Secondary	
3	45559	0.0	2013-01-22	Finn Water	267	FINN WATER	38.058046	-9.418672	Kwa Mzee Pange	
4	49871	500.0	2013-03-27	Bruder	1260	BRUDER	35.006123	-10.950412	Kwa Mzee Turuka	

5 rows × 40 columns

The file "test_set_values" is just like training_set_values, with fewer records (14,850).

```
In [9]: df = pd.read_csv('../data/SubmissionFormat.csv')  
# show row and column counts  
df.shape
```

```
Out[9]: (14850, 2)
```

```
In [10]: # show first few records  
df.head()
```

Out[10]:

	id	status_group
0	50785	predicted label
1	51630	predicted label
2	17168	predicted label
3	45559	predicted label
4	49871	predicted label

The file "SubmissionFormat" is like training_set_labels except the contestant/data scientist has to provide the labels.

Summary of available files

There is a training set consisting of 59,400 records; a set of labels for those 59,400 records; a test set consisting of 14,850 records; and a template for submitting labels for those 14,850 test records to an online contest.

Because we do not have labels for the 14,850 records in the test set, those records unfortunately have no value to us. We will have to carve out training and test sets from just the 59,400 records with known labels.

Create the desired data set

As we drop records from the data set, we'll need to make sure we drop the *same* records from the target labels, so it will be necessary to join those two dataframes together first.

* Special note:

As we'll explore a bit later, many of the "gps_height" values were found to be nonsensical zero values. In a separate notebook we retrieved many of the missing values for this feature that would be relevant to our data set here (i.e. elevations for records whose funder is the tanzanian government). In the next cell, we will reload the training set values, but we'll use an alternate version of this document with the new elevation values as an additional column.

```
In [11]: # load the data from two files  
dfX = pd.read_csv('../data/vals_with_elevation.csv', index_col=0)  
dfy = pd.read_csv('../data/training_set_labels.csv')  
# concatenate the files and rename status feature  
df = pd.concat([dfX, dfy[['status_group']]], axis = 1).rename(columns={'status_group': 'sta  
# show rows and columns  
df.shape
```

Out[11]: (59400, 42)

We're only interested in whether the wells are functional or not, so we can encode the target label accordingly before proceeding with data preparation.

We'll keep a "status" column with just the labels "functional" and "non-functional", because that will make more sense on the legends of figures we'll generate. For statistical purposes, we'll make a parallel feature called "target" and adopt the labeling tradition where the baseline / status quo (functional well) is coded as 0 and the *problem* (non-functional well) is coded as 1.

(We'll need to drop the redundant "status" feature before we get to modeling.)

```
In [12]: # label encode the target feature
df['target'] = df['status'].apply(lambda x: 0 if x == 'functional' else 1)
# relabel the status feature
df['status'] = df['target'].apply(lambda x: 'functional' if x == 0 else 'non-functional')
# show rows and columns
df.shape
```

```
Out[12]: (59400, 43)
```

Data preparation

Duplicates

First we'll address duplicates.

```
In [13]: # check for duplicates (excluding the ids) and show how many duplicates there are
len(df[df.duplicated(subset=df.columns.difference(['id']))])
```

```
Out[13]: 36
```

There appear to be 36 duplicated records. We'll drop them.

```
In [14]: # drop the duplicate records
df.drop(df[df.duplicated(subset=df.columns.difference(['id']))].index, inplace=True)
# show row and column counts
df.shape
```

```
Out[14]: (59364, 43)
```

Trim data to Government-funded wells only

As we'll only be working with records of wells that were funded by the government, we should drop all other records before going any further. Once we've done that, we can drop the funder feature.

We worked on the *funder* column in the separate notebook so that we could focus on just records of wells funded by the government. In that notebook we identified labels such as "tanzania" and "tanza" that we suspected also meant that the Tanzanian Government was the funder.

Having completed that work and edited the funder feature already, we can first drop all records with funder values other than the Tanzanian Government, and then drop the funder column altogether.

```
In [15]: # drop records with funders other than government
df.drop(df[df.funder != 'government of tanzania'].index, inplace=True)
```

```
In [16]: # drop this feature, as its values are all now uniform
df.drop(columns='funder', inplace=True)
# show rows and columns
df.shape
```

```
Out[16]: (9182, 42)
```

Missing values

Now we'll look at which columns have missing values.

```
In [17]: # show columns with missing values and the number of values missing  
df[df.columns[df.isna().any()]].isna().sum()
```

```
Out[17]: installer          4  
subvillage        70  
public_meeting     418  
scheme_management   537  
scheme_name       2757  
permit            41  
dtype: int64
```

For any feature with missing values, our options are:

1. drop the feature
2. preserve the feature by imputing the missing values
3. preserve the feature by dropping the records with missing values

The features we identified here as missing values are all *categorical* features, which means there is little hope for imputing real values unless we choose to use the mode as a default value. Another possible way to impute values for these features is if it made sense to assign the value of "other" to the missing values. Even then, this will only be worthwhile if doing so leaves us with sufficiently *few* unique values for that feature. The reason for this is that in a categorization project such as this one, it simply won't be useful to have a large number of unique values for any single feature.

If, say, 1-4 values comprised something like 70% of the records, then it *might* make sense to put all the rest — including the missing values — into an "other" value pile. But if the value counts are more or less evenly distributed over a greater number of values, then there won't be much hope of imputing, then it won't be worth imputing an "other" value, and we'll have to decide whether to drop the feature or drop the records with missing values.

```
In [18]: # show how many unique values each feature has, for the features that also have missing values  
df[df.columns[df.isna().any()]].nunique()
```

```
Out[18]: installer          129  
subvillage        4000  
public_meeting      2  
scheme_management    10  
scheme_name       815  
permit            2  
dtype: int64
```

Let's start with the worst offender, scheme name. It's missing almost a third of its values, and the values that it has are distributed across 808 unique values. Let's look at counts of its top unique values.

```
In [19]: df.scheme_name.value_counts().head(10)
```

```
Out[19]: Machumba estate pipe line    178  
Government           168  
Borehole             101  
Nasula gravity water supply    93  
Kidia kilemapunda     86  
None                 80  
Una mkolowoni        79  
Olkokola pipe line     79  
Komaka mandaka        73  
Kaisho/Isingiro w       70  
Name: scheme_name, dtype: int64
```

Since we're focusing just on government-funded wells, there does seem to be something potentially interesting about the wells that have Government for their scheme name (which the documentation explains means the government runs those wells). However, there are only 168 of these wells, less than 2% of our current data set. We'll just drop the

```
In [20]: df.drop(columns='scheme_name', inplace=True)
# show row and column counts
df.shape
```

```
Out[20]: (9182, 41)
```

Let's do the same analysis for the subvillage feature since it also has a great number of unique values.

```
In [21]: df.subvillage.value_counts().head(10)
```

```
Out[21]: Majengo      99
Madukani       91
Marurani Kati   59
Marurani Juu     54
Shuleni        47
Kati           39
Mtakuja        39
Muungano       34
Bwawani         32
Sokoni          31
Name: subvillage, dtype: int64
```

This also seems too evenly distributed across too many values to be helpful in modeling. We'll drop it.

```
In [22]: df.drop(columns='subvillage', inplace=True)
# show row and column counts
df.shape
```

```
Out[22]: (9182, 40)
```

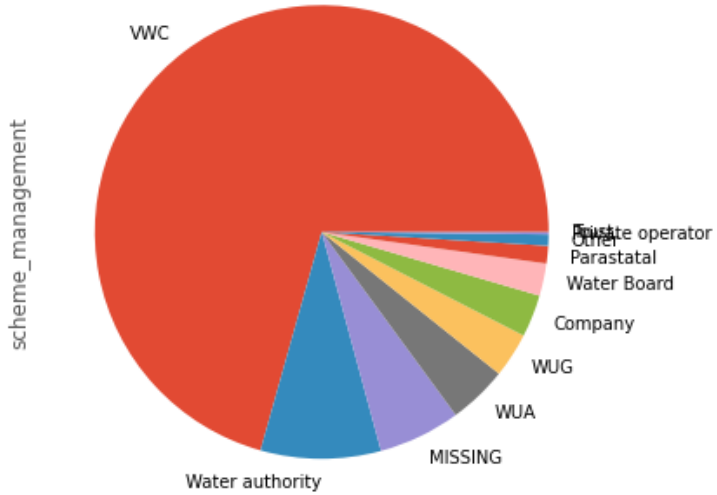
Let's look at scheme management.

```
In [23]: df.scheme_management.value_counts().head(10)
```

```
Out[23]: VWC            6487
Water authority      786
WUA                 386
WUG                 293
Company              277
Water Board           212
Parastatal             15
Other                  73
Private operator       15
Trust                   1
Name: scheme_management, dtype: int64
```

With such a great number concentrated in a few unique values, this looks more promising. Let's look at this as a pie chart, making sure the "missing values" get included as such by relabeling them as "MISSING".

```
In [24]: # relabel missing values
df['scheme_management'].fillna('MISSING', inplace=True)
# plot pie chart of the unique values for scheme_management with highest value counts
df.scheme_management.value_counts().plot.pie(subplots=True, figsize = (11,6), legend=False)
```



Here we see an opportunity to group some values together. Let's replace any value with lower counts than "Other" as "Other", and group the missing values as "Other" as well.

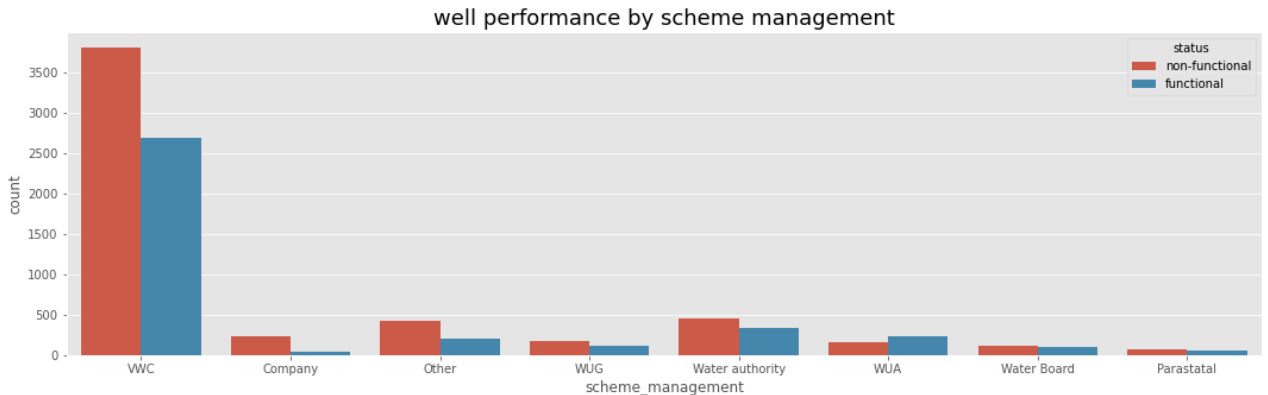
```
In [25]: # create an empty list to be populated by values whose value counts are higher than a certain threshold
other_vals = []

# populate the list
for val in df.scheme_management.value_counts().index:
    if len(df[df.scheme_management == val]) < len(df[df.scheme_management == 'Other']):
        other_vals.append(val)

# reset the values
df['scheme_management'] = df['scheme_management'].apply(lambda x: 'Other' if x in other_vals else x)
# fill any missing values with 'Other'
df['scheme_management'] = df['scheme_management'].apply(lambda x: 'Other' if x == 'MISSING' else x)
```

Let's look at how the wells perform in these categories.

```
In [26]: # set figure
fig, ax = plt.subplots(nrows=1, ncols=1, figsize=(18, 5))
# plot well performance
ax = sns.countplot(x='scheme_management', hue="status", data=df)
# title
ax.set_title('well performance by scheme management', size=18);
```



Let's look at the installer feature.

```
In [27]: df.installer.value_counts().head(15)
```

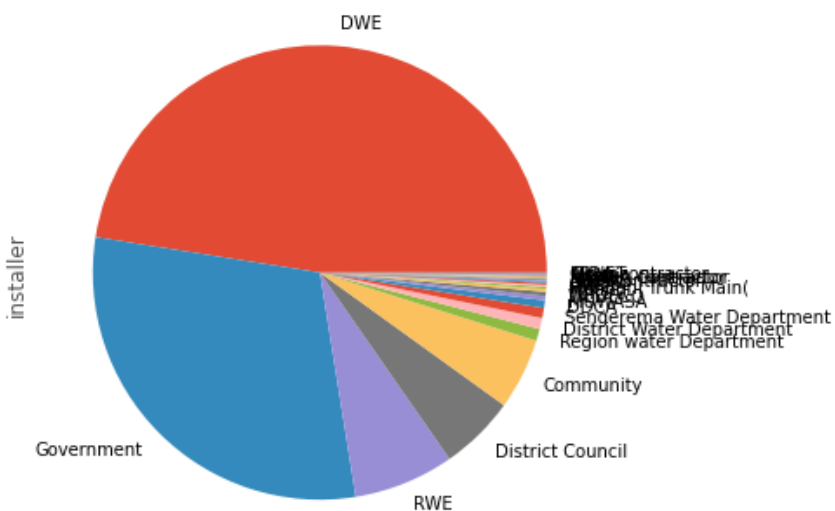
```
Out[27]: DWE                    4309
Government                1600
RWE                      649
District council            295
Commu                     278
Gover                     241
Centr                     162
Community                  159
District Council            150
Central govt                 138
GOVER                     117
Central Government            110
Gove                      91
Central government              90
Region water Department        77
Name: installer, dtype: int64
```

This strongly indicates that some of the labels should be consolidated. We'll make those consolidations before we look at the value counts again.

```
In [28]: df['installer'] = df['installer'].apply(  
    lambda x: 'Government' if x in ['Gover', 'Centr', 'Central govt', 'Central govt', 'Go  
        'Gove', 'Central government', 'Tanzania Government',  
        'Ce', 'Go', 'CENTRAL GOVERNMENT', 'central government  
        'Cebtral Government', 'Tanzanian Government', 'GOVERM  
        'Centra govt', 'Cetral government /RC'] else x)  
df['installer'] = df['installer'].apply(  
    lambda x: 'District Council' if x in ['District council', 'Distri', 'DISTRICT COUNCI  
        'Mbozi District Council', 'Council', 'District  
df['installer'] = df['installer'].apply(  
    lambda x: 'Community' if x in ['Commu', 'RWE /Community', 'RWE/Community', 'RWE/ Comm  
        'RWE Community', 'Communit', 'District Community j'] else x)  
df['installer'] = df['installer'].apply(  
    lambda x: 'DWE' if x in ['Dwe', 'DW'] else x)  
df.installer.value_counts().head(15)
```

Once again, this looks promising. Let's give the missing values a label and visualize with a pie chart again.

```
In [29]: # relabel missing values
df['installer'].fillna('MISSING', inplace=True)
# plot pie chart of 25 unique values for installer with highest value counts
df.installer.value_counts().head(25).plot.pie(subplots=True, figsize = (11,6), legend=False)
```



Here again we can group anything with a value count smaller than that of Community as "other".

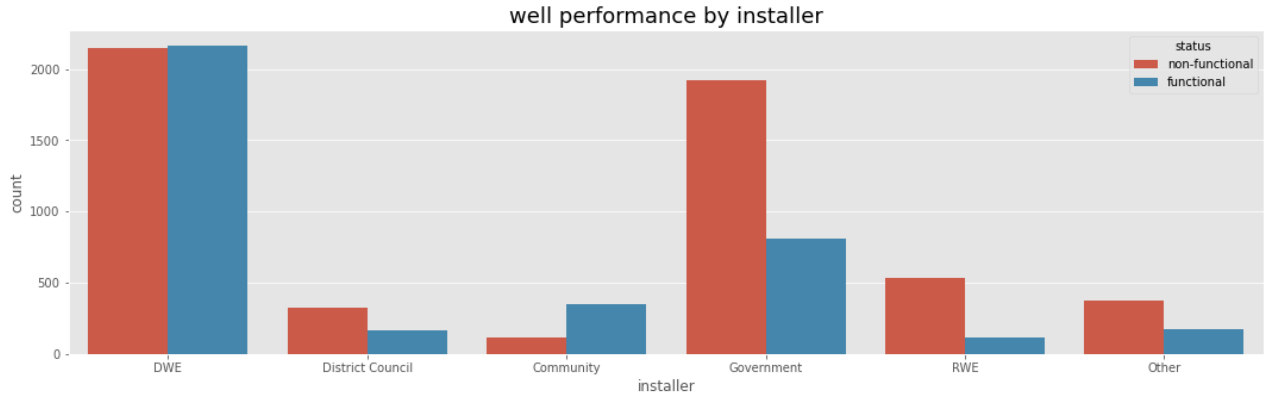
```
In [30]: # create an empty list to be populated by values whose value counts are higher than a ceiling
other_vals = []

# populate the list
for val in df.installer.value_counts().index:
    if len(df[df.installer == val]) < len(df[df.installer == 'Community']):
        other_vals.append(val)

# reset the values
df['installer'] = df['installer'].apply(lambda x: 'Other' if x in other_vals else x)
# fill any missing values with 'Other'
df['installer'] = df['installer'].apply(lambda x: 'Other' if x == 'MISSING' else x)
```

Let's look at how the different installers affect well performance.

```
In [31]: # set figure
fig, ax = plt.subplots(nrows=1, ncols=1, figsize=(18, 5))
# plot well performance
ax = sns.countplot(x='installer', hue="status", data=df)
# title
ax.set_title('well performance by installer', size=18);
```



The remaining features with missing values — public_meeting and permit — have only two unique values each. It could get a little trickier to impute anything here. Let's look at those values, starting with public_meeting.

```
In [32]: df.public_meeting.value_counts()
```

```
Out[32]: True      7929
False     835
Name: public_meeting, dtype: int64
```

```
In [33]: df.permit.value_counts()
```

```
Out[33]: True      6837
False     2304
Name: permit, dtype: int64
```

It's difficult to justify imputing anything here, even an "other" category, so for each of these features we'll have to either drop the feature or drop the records. The missing values for public_meeting are nearly 5% of the data, and we don't even know how helpful this feature will be yet, so it's safe to drop the column, since we the information gain we get from that column is not worth losing 5% of our records.

```
In [34]: df.drop(columns='public_meeting', inplace=True)
# show row and column counts
df.shape
```

```
Out[34]: (9182, 39)
```

There are fewer missing values for the permit feature — only about half of a percent — so it's tempting to just drop those records and keep the feature. It would be nice to know whether this feature might matter.

And because we'll likely run into this question again (of whether a feature will even matter), we'll create a function that can help answer that question for us.

```
In [35]: # create a function that compares functionality success
def compare_vals(feature, vals):
    # loop over all the values in a set of values
    for val in vals:
        # define the numerator as the number of functional records
        num = len(df[(df[feature] == val) & (df['target'] == 0)])
        # define the denominator
        denom = len(df[df[feature] == val])
        # report the percentage
        print(val, "%0.0f%%" % (100 * num / denom))
```

```
In [36]: compare_vals('permit', [True, False])
```

```
True 41%
False 41%
```

This shows that 41% of wells *with* permits are functional, and 41% of wells *without* permits are functional. Now we know for sure that this feature is truly useless and can safely drop it.

```
In [37]: df.drop(columns='permit', inplace=True)
# show row and column counts
df.shape
```

```
Out[37]: (9182, 38)
```

Administrative features

The id, date_recorded, and recorded_by features won't help with any of our analysis. We can safely drop them.

```
In [38]: df.drop(columns=['id', 'date_recorded', 'recorded_by'], inplace=True)
# show row and column counts
df.shape
```

```
Out[38]: (9182, 35)
```

Numerical features with potentially false zeros

We've tackled all the missing values now, but sometimes values of zero are entered when the value is actually missing.

Let's explore the numerical features and find how many zero values each feature has and what percentage of its values are zero.

```
In [39]: for col in df.select_dtypes(include=['number']).columns:
    count = df[col][abs(df[col]) == 0].count()
    pct = round(100*count/(len(df)),2)
    print(col, (19-len(col))*' ', count, (5-len(str(count)))*' ', pct, '%')
```

amount_tsh	7096	77.28 %
gps_height	2599	28.31 %
longitude	234	2.55 %
latitude	0	0.0 %
num_private	9041	98.46 %
region_code	0	0.0 %
district_code	0	0.0 %
population	2841	30.94 %
construction_year	2710	29.51 %
elevation	280	3.05 %
target	3766	41.02 %

For each of these, we'll consider whether the zeros are "real" and what to do about them if they are not.

Region code and district code

First, it would seem that `region_code` and `district_code` are actually *categorical* variables, not numerical. Let's look at a sample of those values and see how many unique values there are for each.

```
In [40]: # show some values for region and district codes
df[['region_code', 'district_code']].head(10)
```

Out[40]:

	region_code	district_code
25	3	4
28	3	1
32	3	4
39	16	1
41	11	3
44	2	2
53	19	6
59	18	2
61	4	1
67	18	1

```
In [41]: # show how many unique values there are for each feature
df[['region_code', 'district_code']].nunique()
```

Out[41]:

region_code	23
district_code	15
dtype:	int64

There's nothing to suggest that region and district codes should be regarded as numerical. We'll convert them to string objects and later apply one-hot encoding to these features.

```
In [42]: # convert region_code to string object
df['region_code'] = df['region_code'].astype('object')
# convert district_code to string object
df['district_code'] = df['district_code'].astype('object')
```

Num_private

Next, nearly 99% of the records have a zero value for the num_private column, and further, the documentation gives no explanation at all of what num_private is. We'll drop this feature.

```
In [43]: # drop num_private
df.drop(columns='num_private', inplace=True)
# show row and column counts
df.shape
```

```
Out[43]: (9182, 34)
```

Construction Year

Let's sample the construction_year values.

```
In [44]: # show some values for construction year
df.construction_year.value_counts().head()
```

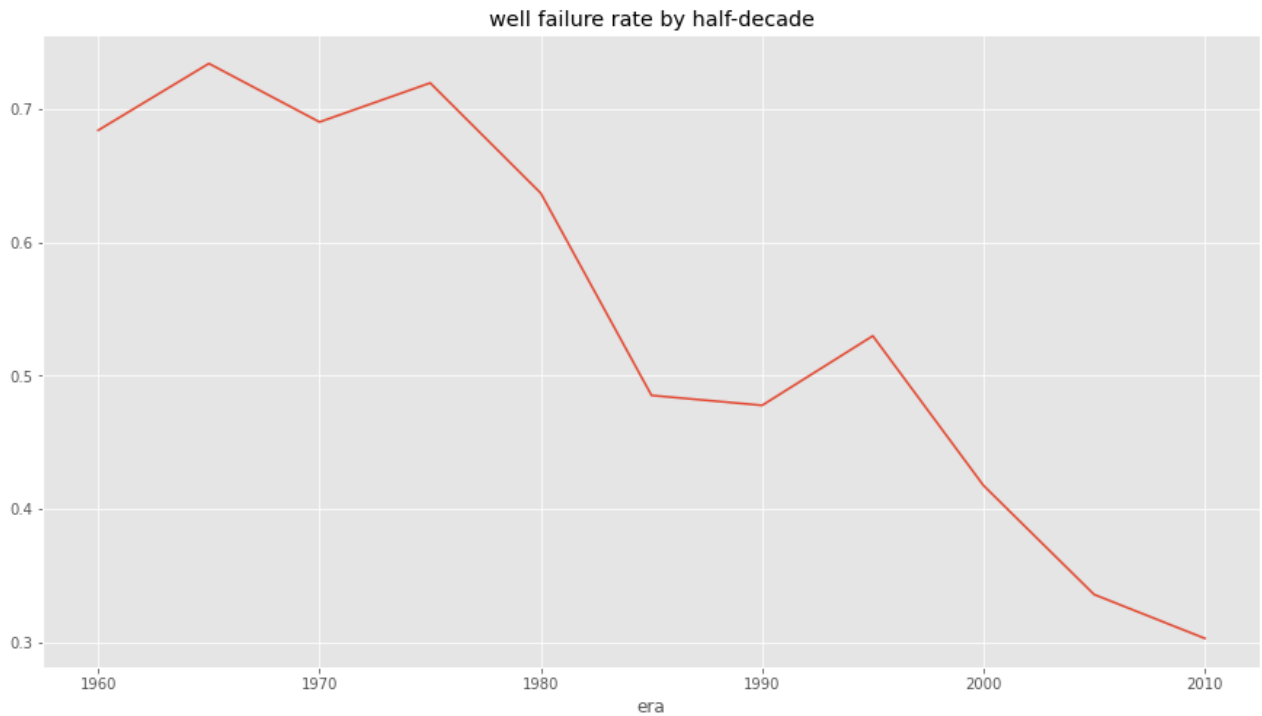
```
Out[44]: 0      2710
1972    514
1974    415
2010    322
1976    279
Name: construction_year, dtype: int64
```

As expected, these zero values make no sense. There's no way to impute the actual values for this feature, and roughly *one third* of the records have missing values in this column, so we'll have to drop that feature.

That's unfortunate, because this feature could potentially have been helpful. But we just can't justify guessing at one third of the values and pretending the result is somehow meaningful.

Before we delete this feature, let's try to learn what we can from the nonzero values. Here we'll make a line graph of performance percentage by half-decade.

```
In [45]: df['era'] = (df['construction_year'] // 5) * 5
df[df.construction_year > 0].groupby('era').mean()['target'].plot(
    figsize=(15,8), title='well failure rate by half-decade');
df.drop(columns='era', inplace=True)
```



It certainly looks like more recently installed wells perform better (fail less).

```
In [46]: # drop construction_year
df.drop(columns=['construction_year'], inplace=True)
# show row and column counts
df.shape
```

```
Out[46]: (9182, 33)
```

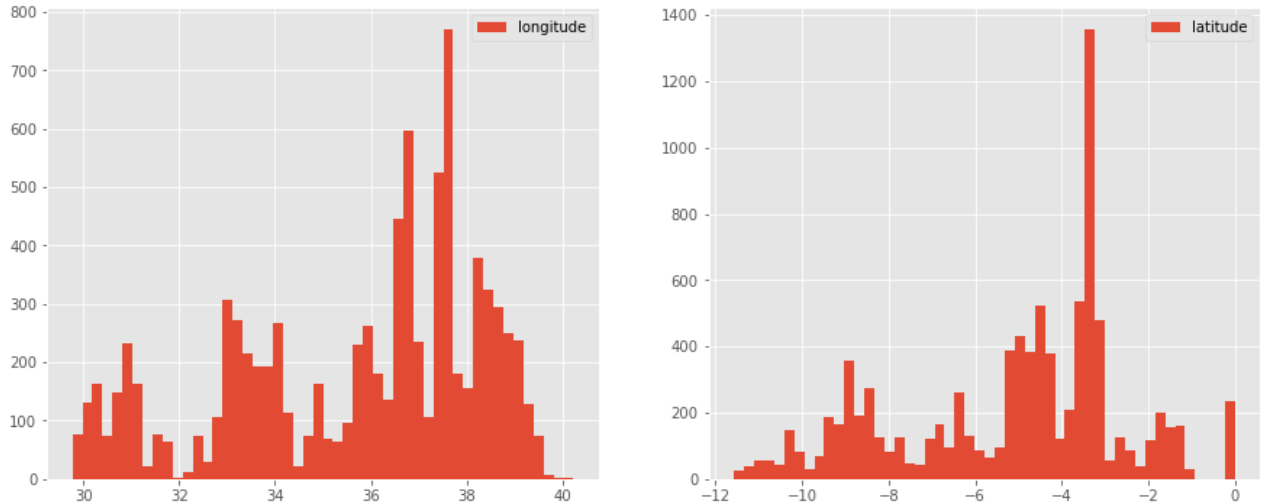
Longitude and latitude

Easily available maps online show that locations within Tanzania's boundaries should have longitude values roughly between 29 and 40, so the zero values for longitude are certainly false.

Latitude values should be roughly between -1 and -11.

Let's look at a histogram of the *non-zero* longitudes and *all* the latitudes to confirm what we have.

```
In [47]: # set figure with two axes over two columns
fig, (ax1, ax2) = plt.subplots(ncols = 2, figsize=(15,6))
# plot histogram of nonzero longitudes
df[df.longitude != 0].longitude.hist(bins=50, ax=ax1, legend=True)
# plot histogram of all latitudes
df.latitude.hist(bins=50, ax=ax2, legend=True);
```



The nonzero longitudes check out fine, but there seem to be some *near-zero* latitudes. Let's take a closer look at these values. Specifically, let's look at the *longitude* values for these near-zero *latitudes*.

```
In [48]: # show longitude values for near-zero latitude records
df[df.latitude > -.2].longitude.describe()
```

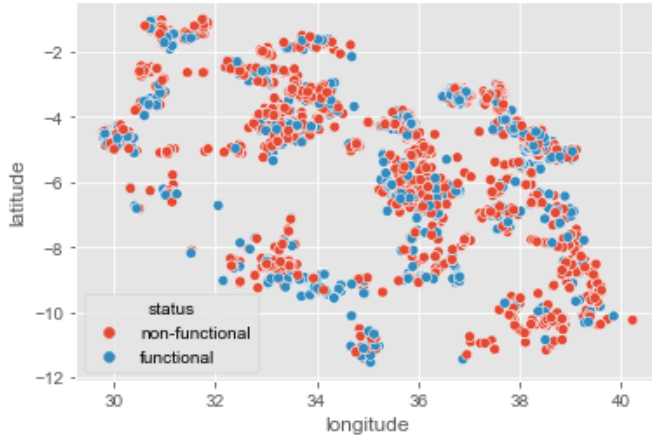
```
Out[48]: count    234.0
mean      0.0
std       0.0
min      0.0
25%      0.0
50%      0.0
75%      0.0
max      0.0
Name: longitude, dtype: float64
```

Just as expected, these near-zero latitude values are for the exact same (234) records that have zero longitude values.

The thing is, we will ultimately drop these two features before modeling anyway, because lat/long are not true numerical features, i.e. 20° of longitude compared to 10° of longitude does not represent *twice as much* of anything. These features were only ever going to help us plot and understand some of the other features. This just means that we'll have to omit those records when we make those plots.

Now that we know which lat/long values we can use, let's generate a map that shows where wells are functional and non-functional

```
In [49]: sns.scatterplot(x="longitude", y="latitude",
                      hue="status",
                      data=df[df.longitude != 0]
                     )
sns.set(rc={'figure.figsize':(10,8)});
```

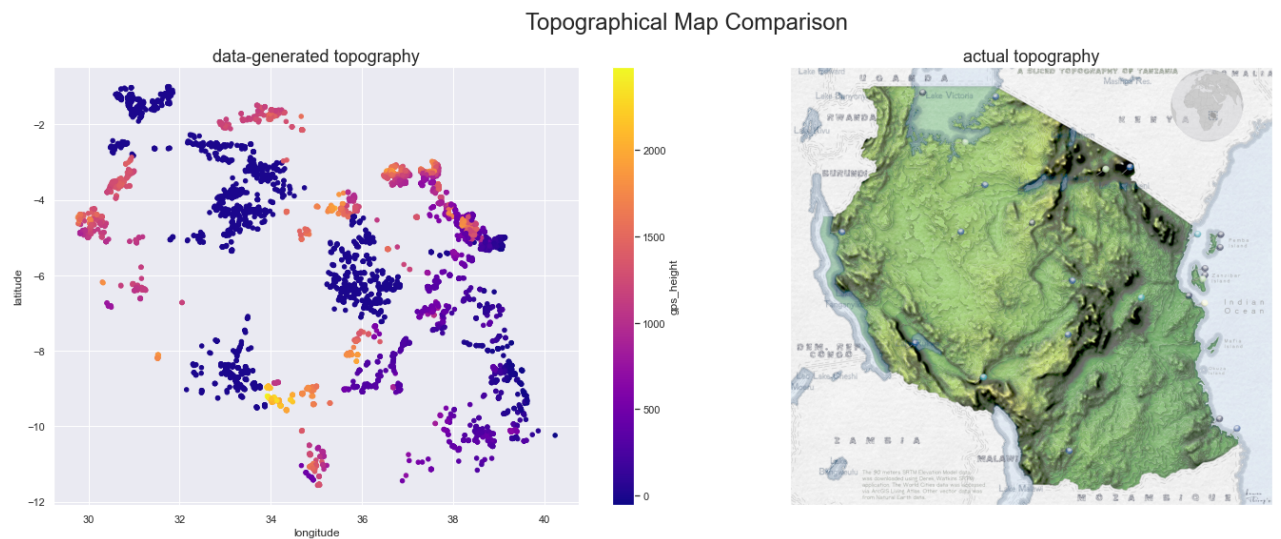


(Non-)functionality doesn't appear to dominate in any general geographical area. Functional wells are regularly interspersed with non-functional wells.

GPS height (altitude)

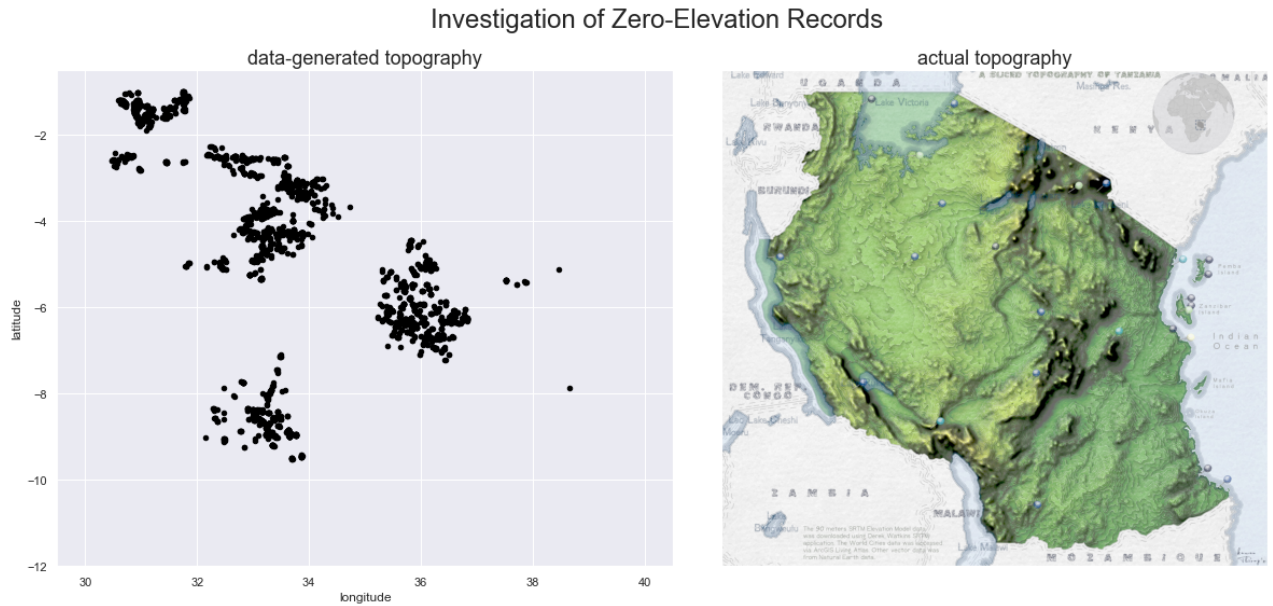
Next let's investigate gps_height. Since Tanzania has a coastline along the Indian Ocean, it's plausible that some locations are in fact at zero elevation. Let's use the (nonzero) lat/long values to explore this.

```
In [50]: # set figure with two axes over two columns
fig, (ax1, ax2) = plt.subplots(ncols = 2, figsize=(20,8))
# plot long/lat for nonzero longs with color gradient for elevation
df[df.longitude > 0].plot.scatter(x='longitude', y='latitude', c='gps_height', cmap='plasma')
# upload an image
im = plt.imread("../images/topo_map.jpeg")
# display the image
im = ax2.imshow(im)
# hide X and Y axes label marks
ax2.xaxis.set_tick_params(labelbottom=False)
ax2.yaxis.set_tick_params(labelleft=False)
# hide X and Y axes tick marks
ax2.set_xticks([])
ax2.set_yticks([])
# titles
fig.suptitle('Topographical Map Comparison', size=24)
ax1.set_title('data-generated topography', size=18)
ax2.set_title('actual topography', size=18)
fig.tight_layout();
```



The data-generated plot on the left indicates that there are lower elevations near the coastline, but there are also many lower elevations in inland areas. A cursory comparison against the actual topographical map on the right suggests those inland areas probably can't be that low, but let's look at a scatterplot of which records are exactly zero.

```
In [51]: # set figure with two axes over two columns
fig, (ax1, ax2) = plt.subplots(ncols = 2, figsize=(17,8))
# plot long/lat of *only* zero elevation records
df[(df.longitude > 0) & (df['gps_height'] == 0)].plot.scatter(
    x='longitude', y='latitude', c = 'black', ax=ax1)
# set the axis ranges to match the ones in the previous plot
ax1.set_xlim([29.5, 40.5])
ax1.set_ylim([-12, -0.5])
# upload an image
im = plt.imread("../images/topo_map.jpeg")
# display the image
im = ax2.imshow(im)
# hide X and Y axes label marks
ax2.xaxis.set_tick_params(labelbottom=False)
ax2.yaxis.set_tick_params(labelleft=False)
# hide X and Y axes tick marks
ax2.set_xticks([])
ax2.set_yticks([])
# title
fig.suptitle('Investigation of Zero-Elevation Records', size=24)
ax1.set_title('data-generated topography', size=18)
ax2.set_title('actual topography', size=18)
fig.tight_layout();
```

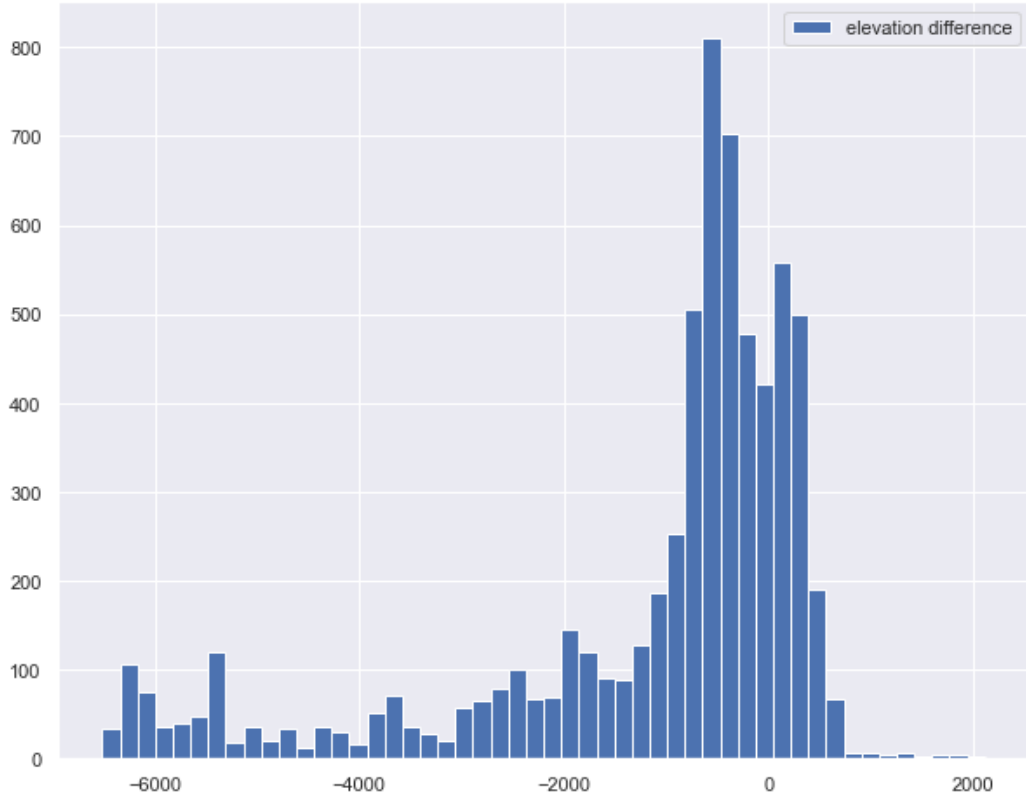


We theorized that some locations could actually have zero elevation, especially if they were along the ocean coastline, but it didn't even turn out that way — the zero-elevation locations are, except for a tiny fraction of them, *all* inland locations. These locations can't possibly (all) have an elevation of (exactly) zero.

In a separate notebook, we were able to implement some api requests to Google to retrieve elevation values for records which we have positional coordinates for. This column with values from Google has the title "elevation".

Unfortunately, a closer look reveals that we won't be able to use either of these features ("gps_height" or "elevation"). If we compare records for which we DO have gps_height values, the elevation values (from Google) are rarely equal. Here is a histogram of the difference between the values:

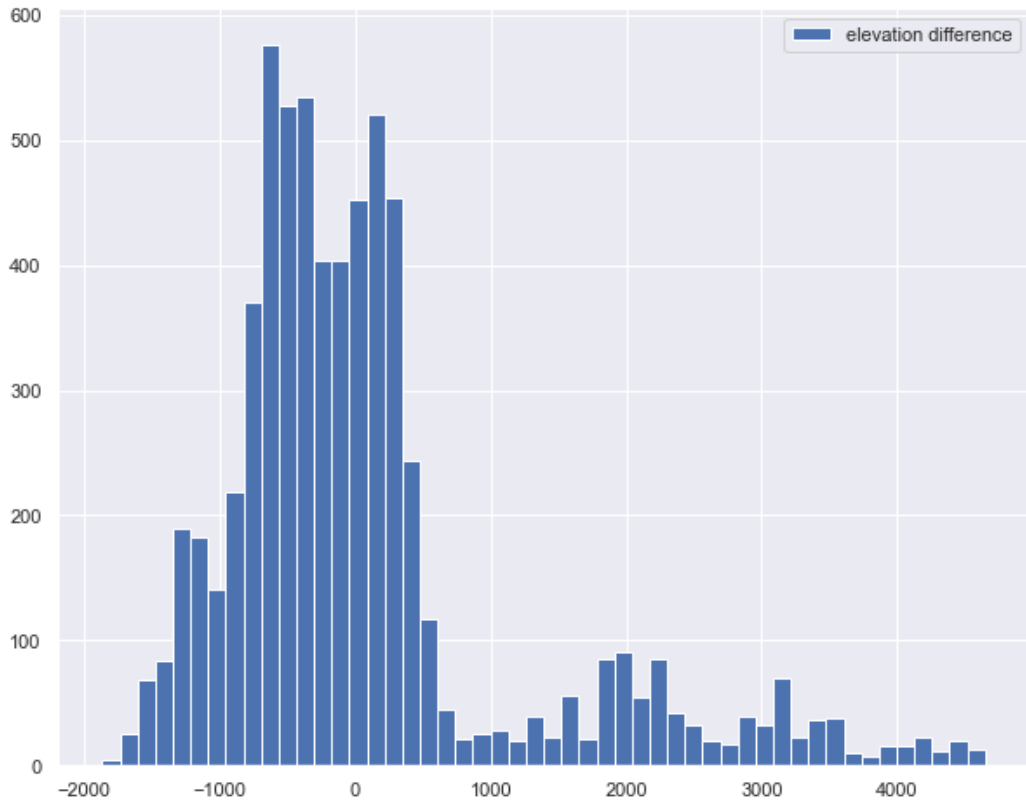
```
In [52]: # create new column of elevation differences
df['elevation difference'] = df['elevation'] - df['gps_height']
# plot a histogram of these differences
df[(df.gps_height != 0)]['elevation difference'].hist(bins=50,legend=True)
# drop the column we created
df.drop(columns='elevation difference', inplace=True);
```



While many differences are close to zero, relatively few are exactly zero, and many differences are way too large.

We could explore whether a sign error were responsible for the differences by making a similar histogram using absolute values of each feature:

```
In [53]: # create new column of differences of absolute values of elevations
df['elevation difference'] = abs(df['elevation']) - abs(df['gps_height'])
# plot a histogram of these differences
df[(df.gps_height != 0)]['elevation difference'].hist(bins=50, legend=True)
# drop the column we created
df.drop(columns='elevation difference', inplace=True);
```



This isn't much better.

We can also explore whether a unit difference (e.g. feet v. meters) is responsible for the differences by examining the ratios of the values. In the case of 3.28 feet = 1 meter, we should expect to see a spike around this value or its reciprocal, 0.30:

```
In [54]: # create new column of (absolute values of) ratios of elevations
df['elevation ratio'] = df.apply(
    lambda x: 0 if x.gps_height == 0 else round(abs(x.elevation / x.gps_height), 2), axis=1
    # show common values for this ratio
)
print(df['elevation ratio'].value_counts().head(10))
# drop the column we created
df.drop(columns='elevation ratio', inplace=True);
```

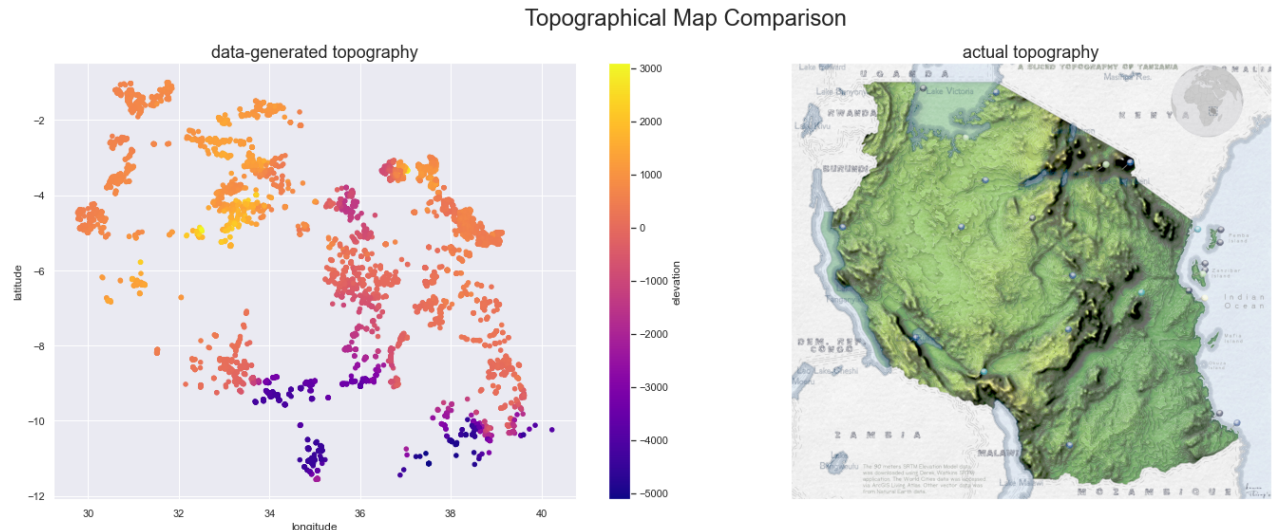
Elevation Ratio	Count
0.00	2624
0.48	91
0.46	89
0.52	80
0.54	80
0.55	75
0.49	71
0.56	70
0.53	70
0.66	69

Name: elevation ratio, dtype: int64

None of the most common values are near enough to either target, 3.28 or 0.30, for us to assume this is a unit error (feet v. meters). Nor do those values seem to cluster around any other ratio that would help us make sense of the discrepancy.

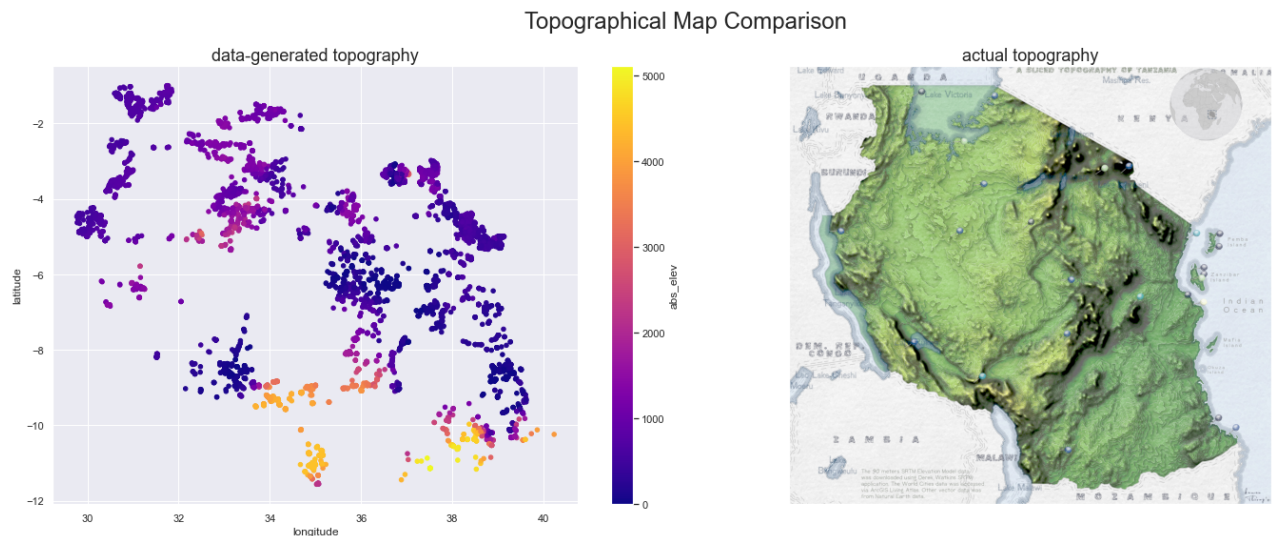
If we simply concluded that the `gps_height` values were to blame, then the elevation values from Google should at least make sense by themselves. Let's make another map and see:

```
In [55]: # set figure with two axes over two columns
fig, (ax1, ax2) = plt.subplots(ncols = 2, figsize=(20,8))
# plot long/lat for nonzero longs with color gradient for elevation
df[df.longitude > 0].plot.scatter(x='longitude', y='latitude', c='elevation', cmap='plasma')
# upload an image
im = plt.imread("../images/topo_map.jpeg")
# display the image
im = ax2.imshow(im)
# hide X and Y axes label marks
ax2.xaxis.set_tick_params(labelbottom=False)
ax2.yaxis.set_tick_params(labelleft=False)
# hide X and Y axes tick marks
ax2.set_xticks([])
ax2.set_yticks([])
# titles
fig.suptitle('Topographical Map Comparison', size=24)
ax1.set_title('data-generated topography', size=18)
ax2.set_title('actual topography', size=18)
fig.tight_layout();
```



Here we see just how many of the values are negative (and too extreme to be real). Our last hope for this feature is if this is only a sign error. Let's try it one more time using absolute values for this feature:

```
In [56]: # create new feature for single use
df['abs_elev'] = abs(df['elevation'])
# set figure with two axes over two columns
fig, (ax1, ax2) = plt.subplots(ncols = 2, figsize=(20,8))
# plot long/lat for nonzero longs with color gradient for elevation
df[df.longitude > 0].plot.scatter(x='longitude', y='latitude', c='abs_elev', cmap='plasma')
# upload an image
im = plt.imread("../images/topo_map.jpeg")
# display the image
im = ax2.imshow(im)
# hide X and Y axes label marks
ax2.xaxis.set_tick_params(labelbottom=False)
ax2.yaxis.set_tick_params(labelleft=False)
# hide X and Y axes tick marks
ax2.set_xticks([])
ax2.set_yticks([])
# titles
fig.suptitle('Topographical Map Comparison', size=24)
ax1.set_title('data-generated topography', size=18)
ax2.set_title('actual topography', size=18)
fig.tight_layout()
# drop the column we created
df.drop(columns='abs_elev', inplace=True);
```



This definitely doesn't look right either. We'll have to drop both the `gps_height` and `elevation` columns.

```
In [57]: # drop gps_height and elevation
df.drop(columns=['gps_height', 'elevation'], inplace=True)
# show row and column counts
df.shape
```

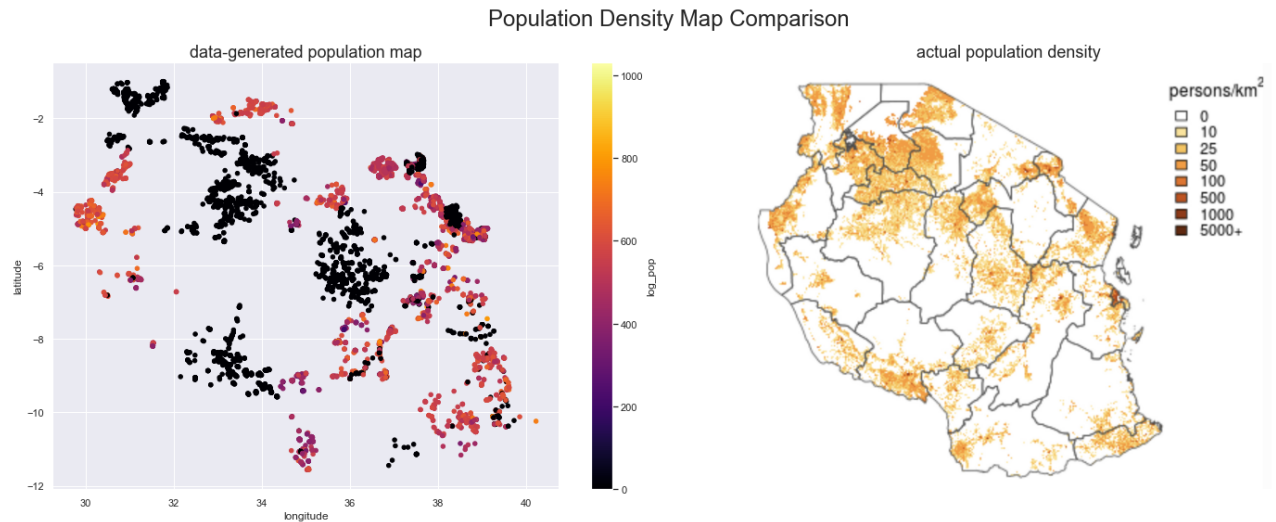
Out[57]: (9182, 31)

Population

The next feature with zero values to explore is population. Let's use another scatterplot map to get a better understanding of this feature.

```
In [58]: # set figure with two axes over two columns
fig, (ax1, ax2) = plt.subplots(ncols = 2, figsize=(20,8))
# create a temporary feature that calculates the logarithm of the population
df['log_pop'] = df.population.apply(lambda x: 0 if x <= 1 else int(np.log(x)*100))
# plot long/lat for nonzero longs with color gradient for population

df[df.longitude > 0].plot.scatter(x='longitude', y='latitude', c='log_pop', cmap='inferno')
# drop the log_pop column that was created
df.drop(columns='log_pop', inplace=True)
# upload an image
im = plt.imread("../images/pop_map.jpeg")
# display the image
im = ax2.imshow(im)
# hide X and Y axes label marks
ax2.xaxis.set_tick_params(labelbottom=False)
ax2.yaxis.set_tick_params(labelleft=False)
# hide X and Y axes tick marks
ax2.set_xticks([])
ax2.set_yticks([])
# title
fig.suptitle('Population Density Map Comparison', size=24)
ax1.set_title('data-generated population map', size=18)
ax2.set_title('actual population density', size=18)
fig.tight_layout();
```



(We've calculated logarithm values of the population records in order to better distinguish the population values from each other.)

On the left, we've plotted populations that are served by the government wells. On the right, for comparison, is a presumably accurate population density map of Tanzania from the UC Davis website (<https://gfc.ucdavis.edu/profiles/rst/tza.html> (<https://gfc.ucdavis.edu/profiles/rst/tza.html>)).

For one thing, we can see that there are water wells in pretty much all the same locations where there are people, which is good! However, our data suggests that there aren't any people at the locations in black. That is *not* consistent with the population density map on the right. As these zero values, again, constitute roughly a third of the records, it looks like we'll have to drop the population feature, too, since it would be too costly to drop those records.

It's not likely that knowing the population in a given area would have factored into explaining whether their wells were functioning, so dropping this feature probably isn't such a great loss.

```
In [59]: # drop population
df.drop(columns='population', inplace=True)
# show row and column counts
df.shape
```

```
Out[59]: (9182, 30)
```

Total static head

The last remaining zero value is for amount_tsh or "total static head". The documentation doesn't do much to clarify this term, but simple search results (e.g. <https://www.rfmacdonald.com/documents/Goulds-Centrifugal%20Pump%20Fundamentals.pdf> (<https://www.rfmacdonald.com/documents/Goulds-Centrifugal%20Pump%20Fundamentals.pdf>)) reveal that the term refers to the vertical distance between the source water level and the level at which the water is accessed. As far as we know, it's quite plausible indeed for this value to be zero, so we'll leave it. In fact, one site (<https://www.linkedin.com/pulse/understanding-chilled-water-pump-head-muneer-ahamed-shaik/> (<https://www.linkedin.com/pulse/understanding-chilled-water-pump-head-muneer-ahamed-shaik/>)) describes "closed loop systems", which by definition have zero total static head.

Let's try to understand just how predictive this feature is. First, let's calculate the functional success rates for zero and nonzero tsh.

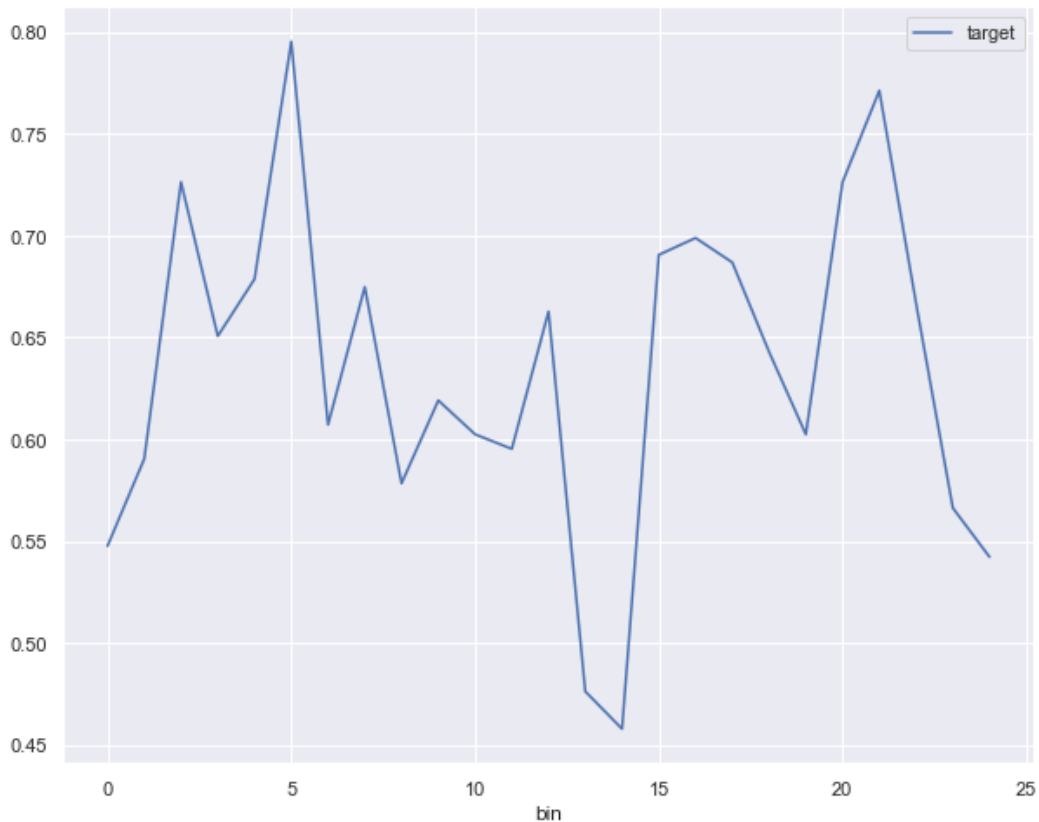
```
In [60]: # the code below is a tweak on the compare_val function
# calculate numerator for zero tsh
num = len(df[(df['amount_tsh'] == 0) & (df['target'] == 0)])
# calculate denominator for zero tsh
denom = len(df[df['amount_tsh'] == 0])
# print as a percentage
print('ZERO tsh:', "%0f%%" % (100 * num / denom))
# calculate numerator for nonzero tsh
num = len(df[(df['amount_tsh'] != 0) & (df['target'] == 0)])
# calculate denominator for nonzero tsh
denom = len(df[df['amount_tsh'] != 0])
# print as a percentage
print('NONZERO tsh:', "%0f%%" % (100 * num / denom))
```

```
ZERO tsh: 34%
NONZERO tsh: 63%
```

So among the supposed "closed loop systems", the well performance is significantly worse than systems with positive values for total static head. (Or are we really just finding that wells perform worse in cases where tsh is *not actually known*?)

The more than 2,000 nonzero tsh values are tricky to explore because they are not distributed uniformly, and there are many repeated values. We'll create and reorder a new dataframe of just these values and then make a "success rate" line graph.

```
In [61]: # create dataframe of only nonzero tsh records and their target values
df_tsh = df[df['amount_tsh'] != 0][['amount_tsh', 'target']].sort_values(
    'amount_tsh', ascending=True).reset_index()
# drop old index column and create new one
df_tsh.drop(columns='index', inplace=True)
df_tsh['bin'] = df_tsh.index
# set number of bins
bins = 25
# recalculate bin number according to percentile (or decile, etc. according to number of
df_tsh['bin'] = df_tsh['bin'].apply(lambda x: bins * x // len(df_tsh))
# merge values to make dataframe of only relevant values
df_tsh_hist = 1 - df_tsh.groupby('bin').mean()[['target']]
df_tsh_hist.plot();
```



This does not shed much light on the predictive nature of total static head.

*Note: In an earlier iteration of this notebook, we ran all the models with and without this feature and found that models performed almost unilaterally better **without** the feature.*

We'll drop this feature.

```
In [62]: # drop total static head
df.drop(columns='amount_tsh', inplace=True)
# show row and column counts
df.shape
```

Out[62]: (9182, 29)

Features with large numbers of unique values

Let's look at other categorical features with too many unique values to factor into our analysis.

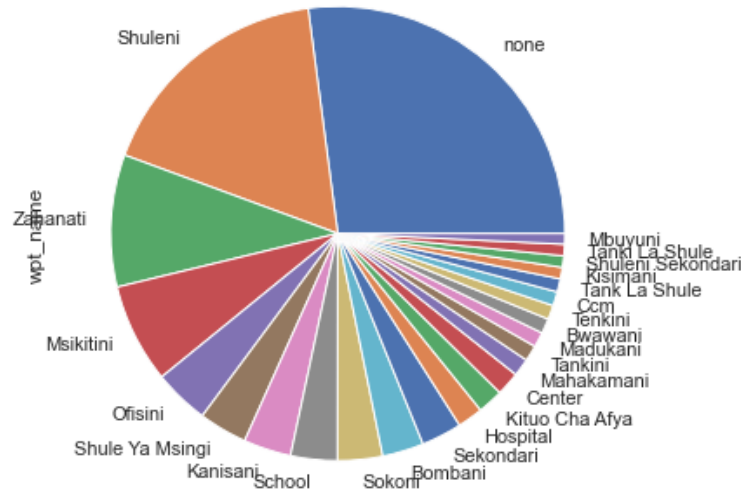
```
In [63]: df.select_dtypes(include='object').loc[  
    :, df.select_dtypes(include='object').apply(lambda x: x.unique().shape[0] > 10).unique()]
```

```
Out[63]: wpt_name          6529  
region             20  
region_code        23  
district_code      15  
lga                81  
ward               901  
extraction_type    17  
extraction_type_group 13  
management         11  
dtype: int64
```

Waterpoint name, Ward, and LGA

We'll investigate wpt_name, ward, and lga using the pie chart method we used earlier.

```
In [64]: # plot pie chart of 25 unique values for wpt_name with highest value counts  
df.wpt_name.value_counts().head(25).plot.pie(subplots=True, figsize = (11,6), legend=False)
```



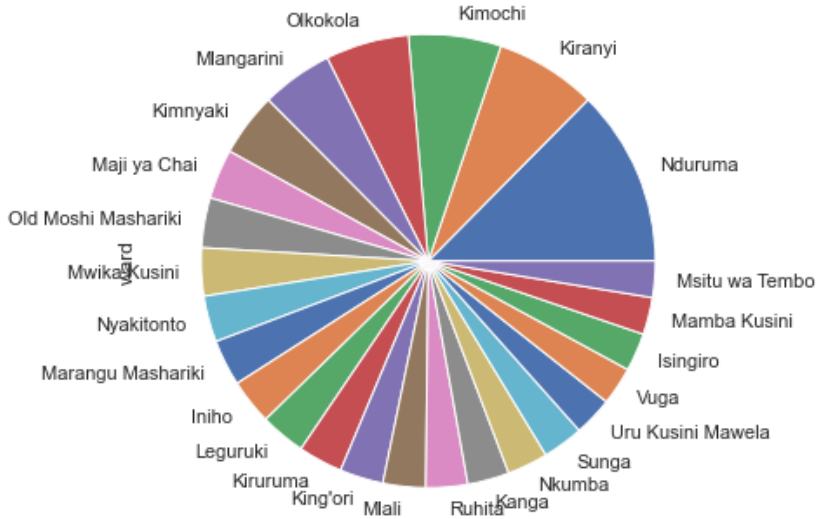
We see that the largest number of these values are actually missing, and the rest are rather evenly distributed, so we can safely drop them.

```
In [65]: df.drop(columns='wpt_name', inplace=True)  
# show row and column counts  
df.shape
```

```
Out[65]: (9182, 28)
```

Here's the pie chart for the ward feature.

```
In [66]: # plot pie chart of 25 unique values for ward with highest value counts
df.ward.value_counts().head(25).plot.pie(subplots=True, figsize = (11,6), legend=False);
```



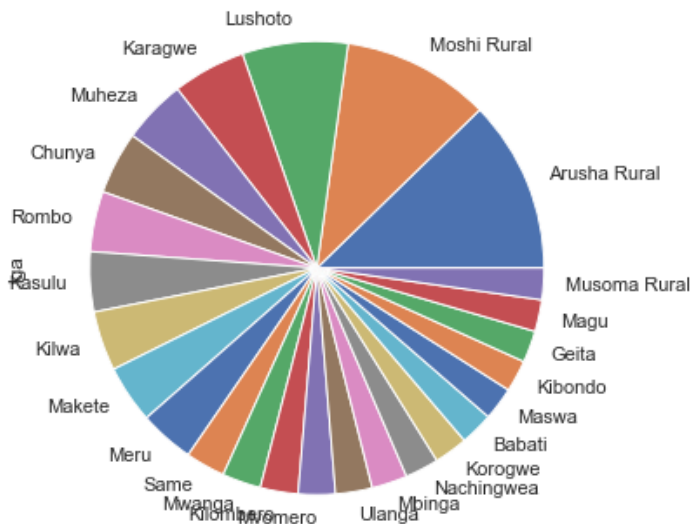
There's no apparent way to group these into anything useable. We'll drop it.

```
In [67]: df.drop(columns='ward', inplace=True)
# show row and column counts
df.shape
```

```
Out[67]: (9182, 27)
```

Here's the pie chart for the lga feature.

```
In [68]: # plot pie chart of 25 unique values for lga with highest value counts
df.lga.value_counts().head(25).plot.pie(subplots=True, figsize = (11,6), legend=False);
```



This will have to go as well.

```
In [69]: df.drop(columns='lga', inplace=True)
# show row and column counts
df.shape
```

```
Out[69]: (9182, 26)
```

The three features we just dropped were all geographical identifiers. It's not clear whether that kind of feature would be useful in our modeling or not.

Geographical identifiers

These features serve as geographical identifiers: basin, region, region_code, and district_code. There is likely to be some redundancy among these features, and we should generally try to understand them better.

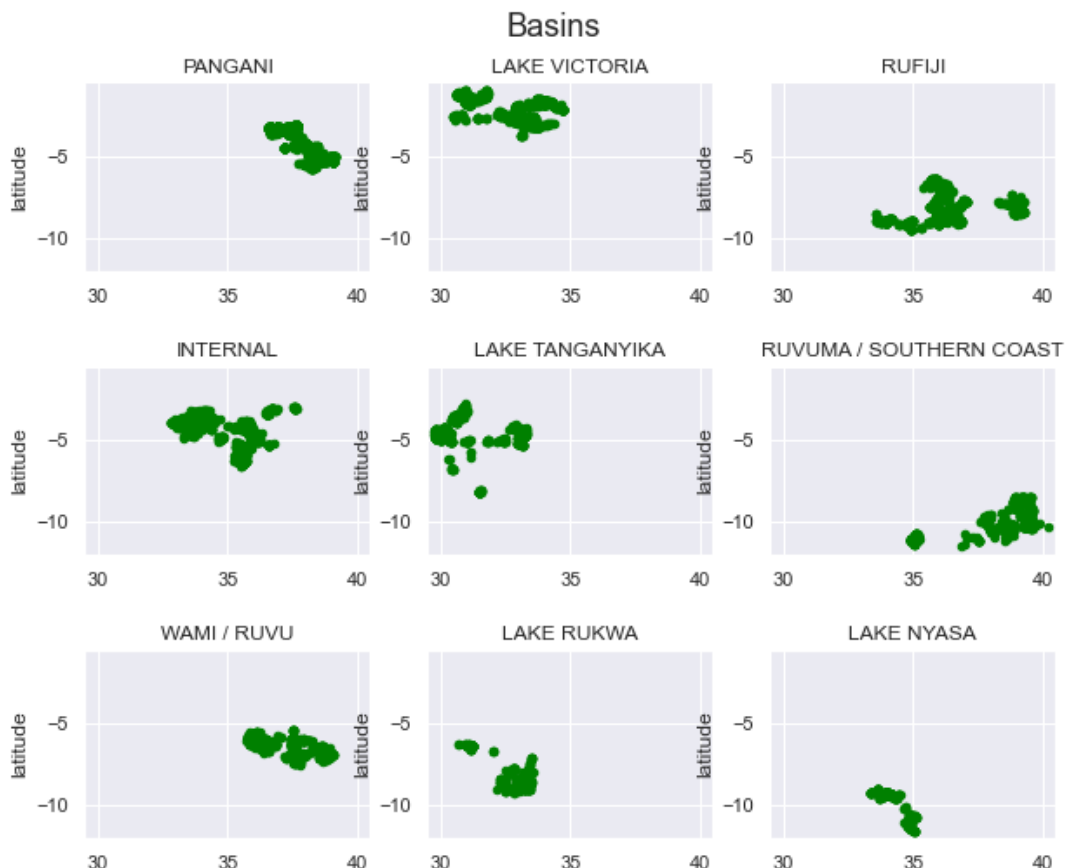
We'll start by plotting all of the basins.

```
In [70]: # define subplot grid
fig, axs = plt.subplots(nrows=3, ncols=3, figsize=(10, 8))
plt.subplots_adjust(hspace=0.5)
fig.suptitle("Basins", fontsize=18, y=0.95)

# loop through basins and axes
for basin, ax in zip(df.basin.value_counts().index, axs.ravel()):
    # filter df for ticker and plot on specified axes
    df[df['basin'] == basin].plot.scatter(x='longitude', y='latitude', c='green', ax=ax)

    # chart formatting
    ax.set_title(basin.upper())
    ax.set_xlabel("")
    ax.set_xlim([29.5, 40.5])
    ax.set_ylim([-12, -0.5])

plt.show();
```

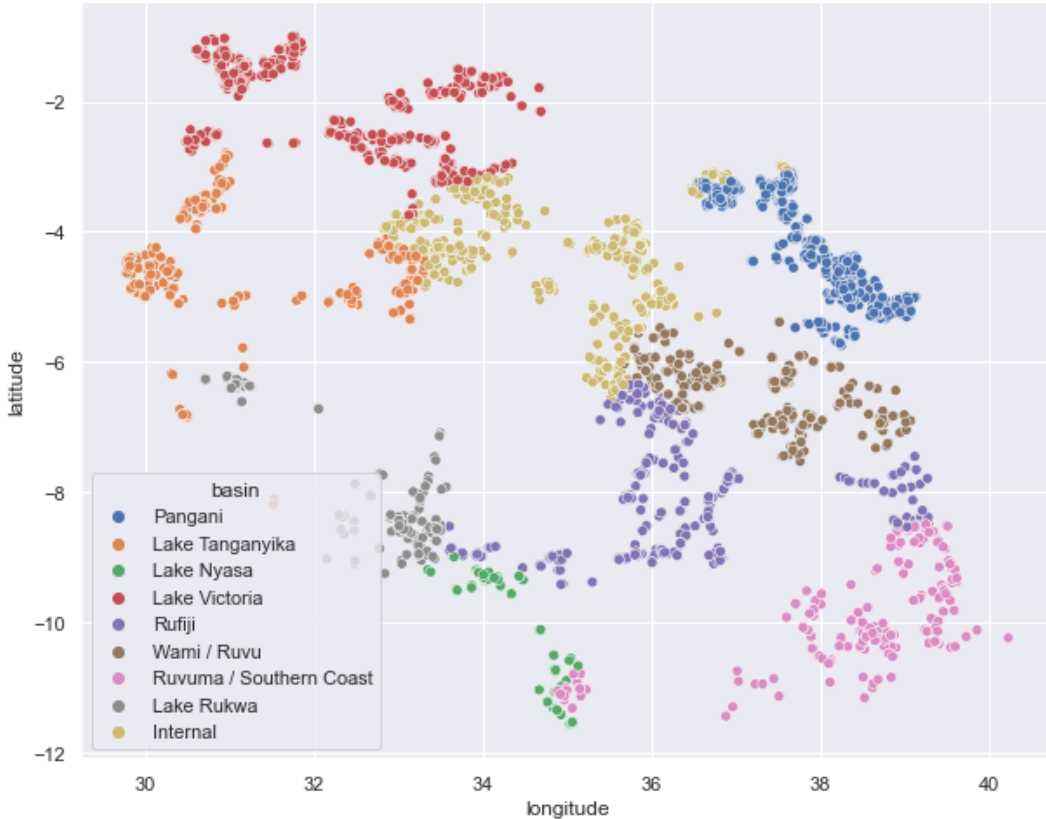


Each basin appears to be roughly contiguous and generally separate from other basins, which looks good.

Let's look at how the basins fit together.

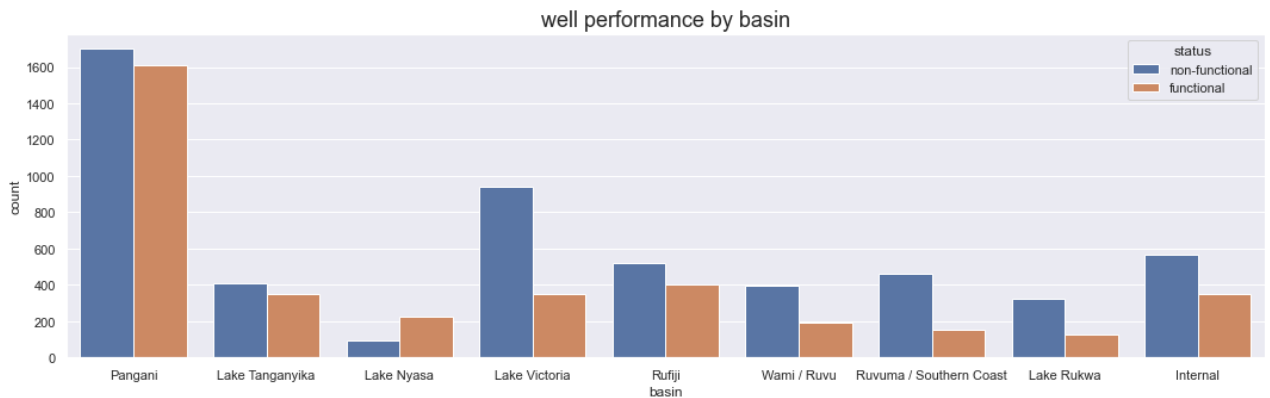
```
In [71]: sns.scatterplot(x="longitude", y="latitude",
                      hue="basin",
                      data=df[df.longitude != 0]
                     )

sns.set(rc={'figure.figsize':(15,12)});
```



Let's take a look at how the wells performed in each basin.

```
In [72]: # set figure
fig, ax = plt.subplots(nrows=1, ncols=1, figsize=(18, 5))
# plot well performance
ax = sns.countplot(x='basin', hue="status", data=df)
# title
ax.set_title('well performance by basin', size=18);
```



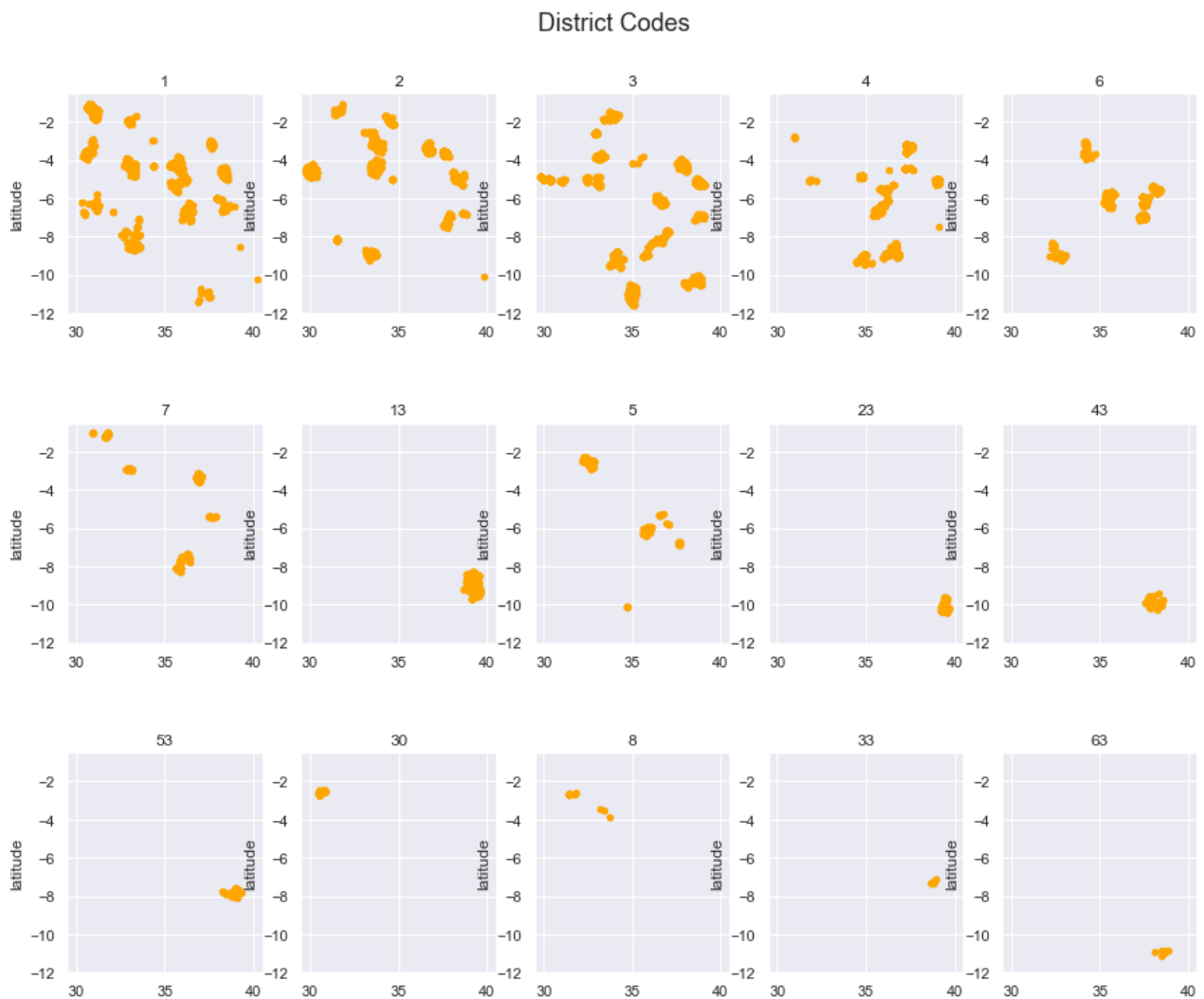
The Pangani basin has the most non-functional wells. The Lake Victoria basin and the Southern Coast basin perform worst by percentage.

```
In [73]: # define subplot grid
fig, axs = plt.subplots(nrows=3, ncols=5, figsize=(15, 12))
plt.subplots_adjust(hspace=0.5)
fig.suptitle("District Codes", fontsize=18, y=0.95)

# loop through basins and axes
for code, ax in zip(df.district_code.value_counts().index, axs.ravel()):
    # filter df for district code and plot on specified axes
    df[df['district_code'] == code].plot.scatter(x='longitude', y='latitude', c='orange')

    # chart formatting
    ax.set_title(str(code).upper())
    ax.set_xlabel("")
    ax.set_xlim([29.5, 40.5])
    ax.set_ylim([-12, -0.5])

plt.show();
```



The first several "districts" seem to overlap a great deal. Since we can't make proper sense of this feature, we'll drop it.

```
In [74]: df.drop(columns='district_code', inplace=True)
# show row and column counts
df.shape
```

```
Out[74]: (9182, 25)
```

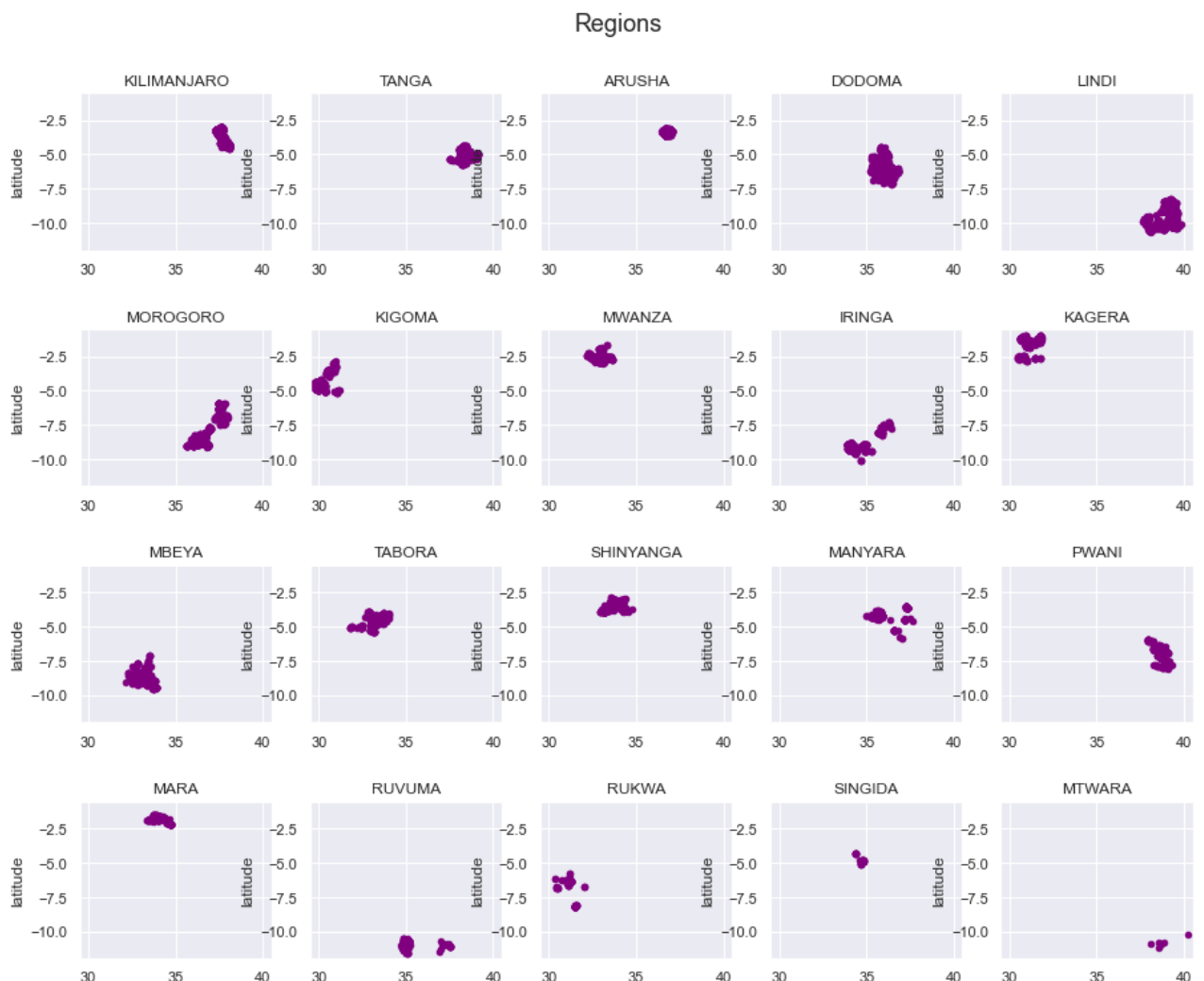
Next, we'll look at the region feature.

```
In [75]: # define subplot grid
fig, axs = plt.subplots(nrows=4, ncols=5, figsize=(15, 12))
plt.subplots_adjust(hspace=0.5)
fig.suptitle("Regions", fontsize=18, y=0.95)

# loop through regions and axes
for region, ax in zip(df.region.value_counts().index, axs.ravel()):
    # filter df for region and plot on specified axes
    df[df['region'] == region].plot.scatter(x='longitude', y='latitude', c='purple', ax=ax)

    # chart formatting
    ax.set_title(region.upper())
    ax.set_xlabel("")
    ax.set_xlim([29.5, 40.5])
    ax.set_ylim([-12, -0.5])

plt.show();
```



These also appear to be contiguous and separate regions.

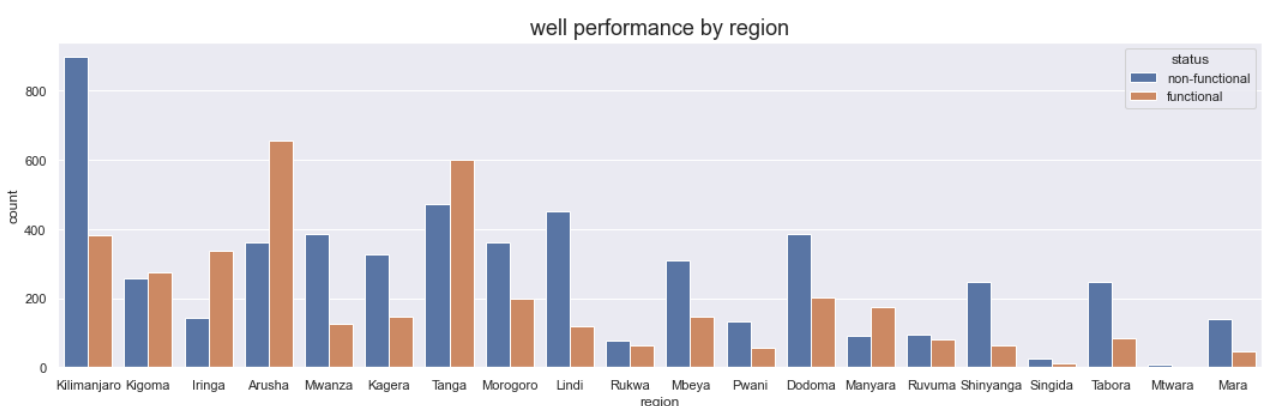
Let's look at how the regions fit together.

```
In [76]: sns.scatterplot(x="longitude", y="latitude",
                      hue="region",
                      data=df[df.longitude != 0]
                     )
sns.set(rc={'figure.figsize':(15,12)});
```



Let's take a look at how the wells performed in each region.

```
In [77]: # set figure
fig, ax = plt.subplots(nrows=1, ncols=1, figsize=(18, 5))
# plot well performance
ax = sns.countplot(x='region', hue="status", data=df)
# title
ax.set_title('well performance by region', size=18);
```



There was a variety of performance levels among the 20 regions.

To recap our geographical identifiers, we already dropped, "ward", "subvillage", "lga", and "wpt_name", and we have now dropped "district code" (and will soon drop region code), leaving just basin and region.

It would seem that "basin" is a natural distinction (a description of natural land features), whereas "region" is man-made (imposed by governing bodies). It may be interesting to note which of these ultimately proves more meaningful.

Possibly redundant features

Several features seem to be related, such as waterpoint_type and waterpoint_type_group. In such cases, we would generally prefer to keep the one with fewer unique values and drop the other(s). Let's look at value counts for groups of features that seem to be related.

To help in this part of the exploration, we'll often employ the `compare_vals` function we created much earlier to help us determine how certain values of a feature affect the target feature, i.e. whether those values serve a useful distinction or they should be lumped together.

region / code

```
In [78]: df[['region_code', 'region']].value_counts()
```

```
Out[78]: region_code    region
3             Kilimanjaro    1280
4              Tanga        1072
2              Arusha        1017
1              Dodoma         589
5            Morogoro         562
16             Kigoma         533
19             Mwanza         513
11             Iringa         478
18             Kagera         474
12             Mbeya          457
80             Lindi          418
14             Tabora         332
17            Shinyanga        309
21            Manyara         265
20              Mara          185
10            Ruvuma          176
8              Lindi          151
15            Rukwa           138
6              Pwani          130
60            Pwani           59
13            Singida         34
90            Mtwarra          7
18             Lindi           2
99            Mtwarra          1
dtype: int64
```

There are only a very few repetitions of regions with different code numbers here. We'll keep "region", since there are fewer of them, and we'll drop "region code".

```
In [79]: df.drop(columns='region_code', inplace=True)
# show row and column counts
df.shape
```

```
Out[79]: (9182, 24)
```

Extraction type / group / class

```
In [80]: df[['extraction_type', 'extraction_type_group']].value_counts()
```

```
Out[80]: extraction_type      extraction_type_group
gravity                      gravity                  5591
other                        other                   858
mono                         mono                   666
submersible                   submersible             568
nira/tanira                  nira/tanira            490
ksb                          submersible             301
swn 80                       swn 80                 282
india mark ii                india mark ii          190
afridev                      afridev                130
other - rope pump             rope pump               28
other - swn 81                other handpump         28
climax                       other motorpump        21
windmill                      wind-powered           14
india mark iii                india mark iii          5
cemo                          other motorpump        5
other - play pump              other handpump         4
other - mkulima/shinyanga    other handpump         1
dtype: int64
```

The only significant consolidation done here is the types "submersible" and "ksb" get consolidated into the group "submersible". Let's see how distinct these types are.

```
In [81]: compare_vals('extraction_type', ['submersible', 'ksb'])
```

```
submersible 48%
ksb 6%
```

This shows that a far greater percentage "submersible" waterpoints are functional than are the waterpoints of "ksb" extraction type. Our choice (to drop one of two similar columns) involves consolidating these two values into one category or preserving the distinction. Since the percentage difference here seems so significant, it may make the most sense to preserve this distinction. As for the remaining values, we can just consolidate everything with a group size smaller than "afridev" into the "other" group.

```
In [82]: # reassign values in the "submersible" group
df['extraction_type_group'] = df.apply(
    lambda x: 'ksb' if x.extraction_type == 'ksb' else x.extraction_type_group, axis=1)

# create an empty list to be populated by values whose value counts are higher than a certain threshold
other_vals = []

# populate the list
for val in df.extraction_type_group.value_counts().index:
    if len(df[df.extraction_type_group == val]) < len(df[df.extraction_type_group == 'afridev']):
        other_vals.append(val)

# reset the values
df['extraction_type_group'] = df['extraction_type_group'].apply(lambda x: 'other' if x in other_vals else x)
```

Having edited the extraction group, we can drop the extraction type feature.

```
In [83]: df.drop(columns='extraction_type', inplace=True)
# show row and column counts
df.shape
```

```
Out[83]: (9182, 23)
```

Now we need to further consider which feature to keep between extraction type "group" and "class". We'll look at the value counts.

```
In [84]: df[['extraction_type_group', 'extraction_type_class']].value_counts()
```

```
Out[84]: extraction_type_group  extraction_type_class
gravity                 gravity          5591
other                  other           858
mono                   motorpump        666
submersible             submersible      568
nira/tanira             handpump         490
ksb                     submersible      301
swn 80                  handpump         282
india mark ii            handpump         190
afridev                 handpump         130
other                   handpump         38
                           rope pump          28
                           motorpump         26
                           wind-powered       14
dtype: int64
```

We may explore this more in depth later or change our minds during modeling, but for now we'll keep the feature with fewer unique values and drop the other.

```
In [85]: df.drop(columns='extraction_type_group', inplace=True)
# show row and column counts
df.shape
```

```
Out[85]: (9182, 22)
```

Management / group

```
In [86]: df[['management', 'management_group']].value_counts()
```

```
Out[86]: management      management_group
vwc              user-group        7242
wua              user-group        380
water board     user-group        362
wug              user-group        322
company          commercial       258
water authority commercial       182
other             other            171
parastatal       parastatal       129
private operator commercial       69
unknown           unknown           65
trust              commercial        2
dtype: int64
```

Based on the value counts, it seems best to keep the different user-groups but consolidate the rest into "commercial" and "other".

```
In [87]: # reassign values
df['management'] = df.apply(
    lambda x: 'commercial' if x.management_group == 'commercial' else x.management, axis=1)

# create an empty list to be populated by values whose value counts are higher than a certain threshold
other_vals = []

# populate the list
for val in df.management.value_counts().index:
    if len(df[df.management == val]) < len(df[df.management == 'wug']):
        other_vals.append(val)

# reset the values
df['management'] = df['management'].apply(lambda x: 'other' if x in other_vals else x)
```

Then we can drop the other feature.

```
In [88]: df.drop(columns='management_group', inplace=True)
# show row and column counts
df.shape
```

```
Out[88]: (9182, 21)
```

payment / type

```
In [89]: df[['payment', 'payment_type']].value_counts()
```

```
Out[89]: payment           payment_type
never pay            never pay      4766
unknown              unknown       1220
pay monthly          monthly       1216
pay per bucket       per bucket   1187
pay when scheme fails on failure 366
pay annually         annually     296
other                other        131
dtype: int64
```

These are in a precise one-to-one correspondence (they're just named differently), so one of them can be dropped without question. Also, we might as well merge "unknown" with "other".

```
In [90]: df.drop(columns='payment_type', inplace=True)
df['payment'] = df['payment'].apply(lambda x: 'other' if x == 'unknown' else x)
# show row and column counts
df.shape
```

```
Out[90]: (9182, 20)
```

Water quality / group

```
In [91]: df[['water_quality', 'quality_group']].value_counts()
```

```
Out[91]: water_quality    quality_group
soft            good           8117
salty           salty           637
unknown         unknown          298
fluoride        fluoride          44
milky           milky            36
coloured        colored           27
salty abandoned salty             19
fluoride abandoned fluoride            4
dtype: int64
```

Most of the values here seem too small to matter, so we'll keep the feature that's more consolidated and drop "water quality".

```
In [92]: df.drop(columns='water_quality', inplace=True)
# show row and column counts
df.shape
```

```
Out[92]: (9182, 19)
```

Water quantity / group

```
In [93]: df[['quantity', 'quantity_group']].value_counts()
```

```
Out[93]: quantity    quantity_group
enough      enough           4738
insufficient insufficient       3050
dry          dry              983
seasonal    seasonal          316
unknown     unknown            95
dtype: int64
```

This is another one-to-one match, so we can just drop one of them.

```
In [94]: df.drop(columns='quantity_group', inplace=True)
# show row and column counts
df.shape
```

```
Out[94]: (9182, 18)
```

source / type / class

```
In [95]: df[['source', 'source_type']].value_counts()
```

```
Out[95]: source          source_type
          spring           spring        4028
          machine dbh      borehole     1494
          river            river/lake   1399
          shallow well    shallow well  1365
          lake             river/lake   321
          rainwater harvesting rainwater harvesting  252
          dam              dam         243
          hand dtw         borehole     51
          other            other        18
          unknown          other        11
          dtype: int64
```

The only distinctions between source and source_type are that "source" recognizes two values for each of borehole, river/lake, and other. In every case, at least one of the value counts is small enough that there doesn't seem to be any harm in consolidating, so we'll keep "source type" and drop "source".

```
In [96]: df.drop(columns='source', inplace=True)
# show row and column counts
df.shape
```

```
Out[96]: (9182, 17)
```

Let's consider the distinctions between source type and source class

```
In [97]: df[['source_type', 'source_class']].value_counts()
```

```
Out[97]: source_type          source_class
          spring            groundwater    4028
          river/lake         surface       1720
          borehole           groundwater   1545
          shallow well      groundwater   1365
          rainwater harvesting surface     252
          dam               surface       243
          other              unknown      29
          dtype: int64
```

We'll use our function to make some comparisons.

```
In [98]: compare_vals('source_type', ['spring', 'borehole', 'shallow well'])
```

```
spring 50%
borehole 32%
shallow well 36%
```

```
In [99]: compare_vals('source_type', ['river/lake', 'rainwater harvesting', 'dam'])
```

```
river/lake 33%
rainwater harvesting 59%
dam 18%
```

```
In [100]: compare_vals('source_class', ['groundwater', 'surface'])
```

```
groundwater 43%
surface 34%
```

These all seem like interesting enough differences for the numbers of values involved. It's difficult to tell which one might be of more use to us.

At this point, we'll choose to keep the feature with fewer unique values and drop the other, but we could possibly revisit this later depending on how modeling goes.

```
In [101]: df.drop(columns='source_type', inplace=True)
# show row and column counts
df.shape
```

```
Out[101]: (9182, 16)
```

Waterpoint type / group

```
In [102]: df[['waterpoint_type', 'waterpoint_type_group']].value_counts()
```

```
Out[102]: waterpoint_type           waterpoint_type_group
communal standpipe      communal standpipe      5312
communal standpipe multiple  communal standpipe      1417
hand pump                  hand pump              1195
other                      other                  1150
improved spring            improved spring          77
cattle trough               cattle trough          30
dam                         dam                   1
dtype: int64
```

The only difference here is that `waterpoint_type` distinguishes between two types of "communal standpipe". Let's see whether that makes any difference. The function created earlier will tell us whether this distinction matters and should be kept.

```
In [103]: compare_vals('waterpoint_type', ['communal standpipe', 'communal standpipe multiple'])
```

```
communal standpipe 49%
communal standpipe multiple 23%
```

This shows the difference in functional percentage of wells that have "communal standpipe" versus "communal standpipe multiple". This seems like an important difference, so we'll keep the feature that makes that distinction and drop the other feature.

We'll also add the single "dam" value to the "other" category.

```
In [104]: df.drop(columns='waterpoint_type_group', inplace=True)
df['waterpoint_type'] = df['waterpoint_type'].apply(lambda x: 'other' if x == 'dam' else
# show row and column counts
df.shape
```

```
Out[104]: (9182, 15)
```

This should conclude the data preparation, with the exception of dropping latitude and longitude. We won't use latitude and longitude in modeling, but we may want to look at them later. We'll keep them in this dataframe but remember to drop them in the next section. We also must remember to drop the redundant "status" feature, as it is covered numerically by the "target" feature.

Final data preparation steps

Train and test sets

```
In [105]: # split the data into target and predictors
y = df['target']
X = df.drop(columns='target')
# split the data into training and test sets
X_train, X_test, y_train, y_test = train_test_split(X, y, random_state=SEED)
```

One-hot encoding

It is a necessary step to one-hot encode all of the categorical features.

```
In [106]: # one-hot encode the data
X_train = pd.get_dummies(X_train.drop(columns='status'))
X_test = pd.get_dummies(X_test.drop(columns='status'))

# drop longitude and latitude features
X_train.drop(columns=['longitude', 'latitude'], inplace=True)
X_test.drop(columns=['longitude', 'latitude'], inplace=True)
# select one "neutral" predictor to delete from each feature to eliminate redundancy in one-hot encoding
X_train.drop(columns = [
    'installer_Other', 'basin_Lake Nyasa', 'region_Mt悟ara', 'scheme_management_Other', 'education_level_other',
    'management_other', 'payment_other', 'quality_group_unknown', 'quantity_unknown', 'sector_agriculture',
    'waterpoint_type_other'
], axis = 1, inplace = True)
X_test.drop(columns = [
    'installer_Other', 'basin_Lake Nyasa', 'region_Mt悟ara', 'scheme_management_Other', 'education_level_other',
    'management_other', 'payment_other', 'quality_group_unknown', 'quantity_unknown', 'sector_agriculture',
    'waterpoint_type_other'
], axis = 1, inplace = True)
# show row and column counts
X_train.shape
```

```
Out[106]: (6886, 71)
```

```
In [107]: X_test.shape
```

```
Out[107]: (2296, 71)
```

Summary of prepared data

We now have just 11 predictive features (all of them categorical) and 1 target variable for about 9,000 records. One-hot encoding has expanded the 11 predictor columns into about 70 columns.

Modeling

As discussed in our business understanding, we are particularly interested in the **recall** statistic for non-functional wells. To be clear, this recall statistic will show the number of true positives (correctly identified non-functional wells, coded as "1" in the target feature) divided by the (total) number of (actual) non-functional wells. The reason for this interest specifically in recall, again, is to minimize the error of identifying a well as functional when it is not, thus depriving its users of available water.

We'll create a handy function that will show both this recall statistic and the overall accuracy for each model. The

```
In [108]: # function that returns the recall and accuracy statistics as described above
def summary(y_test, y_pred):
    # calculate true positives
    true_pos = confusion_matrix(y_test, y_pred)[1][1] # CORRECTED FROM [0][0] TO [1][1]
    # calculate all actual positives
    all_pos = confusion_matrix(y_test, y_pred)[1].sum() # CORRECTED FROM 0 TO 1
    # calculate true negatives
    true_neg = confusion_matrix(y_test, y_pred)[0][0] # CORRECTED FROM [1][1] TO [0][0]
    # calculate all values
    all_all = confusion_matrix(y_test, y_pred).sum() # CORRECT ALL ALONG
    # return relevant statistics
    return round(true_pos/all_pos,3), round(((true_pos + true_neg)/all_all),3)
```

```
In [109]: # function that reports the recall and accuracy statistics for the train and test sets
def report(y_train, y_train_pred, y_test, y_test_pred):
    y = [[y_train, y_train_pred], [y_test, y_test_pred]]
    title = ['TRAIN SET', 'TEST SET']
    for i in range(2):
        # calculate true positives
        true_pos = confusion_matrix(y[i][0], y[i][1])[1][1]
        # calculate all actual positives
        all_pos = confusion_matrix(y[i][0], y[i][1])[1].sum()
        # calculate true negatives
        true_neg = confusion_matrix(y[i][0], y[i][1])[0][0]
        # calculate all values
        all_all = confusion_matrix(y[i][0], y[i][1]).sum()
        print(title[i], 'recall accuracy')
        print(' ', round(true_pos/all_pos,3), ' ', round(((true_pos + true_neg)/all_all),3))
        if i == 0:
            print('')
```

Before we run any models, let's look at the test data. We'll make a fake prediction that *all* wells are non-functional and run the above function on it.

```
In [110]: # create array with space for all labels
y_pred=np.empty(len(X_test))
# fill this array with all 1s, indicating that we're predicting all wells to be non-functional
y_pred.fill(1)

# show relevant statistics
print('recall: ', summary(y_test,y_pred)[0])
print('accuracy: ', summary(y_test,y_pred)[1])
```

recall: 1.0
accuracy: 0.591

Of course the recall is perfect — we correctly recalled all the non-functional wells, because this "model" predicted all wells would be non-functional.

The accuracy here tells us what percentage of the wells in the test set are actually non-functional.

This is the "dumbest" possible model. All future models had better have better accuracy.

Getting into the weeds of the mathematics just a little: The accuracy is a weighted average of two recall statistics, the recall of non-functional wells and the recall of functional wells. It stands to reason that *one* of those recall values will be higher than the accuracy statistic, and the other will be lower. So we should be able to tweak our models such that the preferred recall statistic is higher than the accuracy statistic.

Logistic Regression

This model uses basic logistic regression.

```
In [111]: # instantiate a LogisticRegression
logreg = LogisticRegression(fit_intercept=False, C=1e12, solver='liblinear', random_state=SEED)
# fit to the training data
logreg.fit(X_train, y_train)
# make predictions for test data
y_train_pred = logreg.predict(X_train)
y_test_pred = logreg.predict(X_test)
# show relevant statistics
report(y_train, y_train_pred, y_test, y_test_pred)
```

TRAIN SET recall accuracy
 0.776 0.757

TEST SET recall accuracy
 0.772 0.75

We'll consider this our baseline model. We're off to a good start in that the preferred recall statistic is higher than the accuracy statistic.

Decision Trees

Here we'll use a decision tree classifier and set the max depth to 5.

```
In [112]: # instantiate a DecisionTreeClassifier
dt_clf = DecisionTreeClassifier(criterion='entropy', max_depth=5, random_state=SEED)
# fit to the training data
dt_clf.fit(X_train, y_train)
# make predictions for the test data
y_train_pred = dt_clf.predict(X_train)
y_test_pred = dt_clf.predict(X_test)
# show relevant statistics
report(y_train, y_train_pred, y_test, y_test_pred)
```

TRAIN SET recall accuracy
 0.76 0.721

TEST SET recall accuracy
 0.763 0.72

This model has slightly diminished recall and more significantly diminished accuracy.

We can (hopefully) make the decision tree model better by hypertuning the max depth.

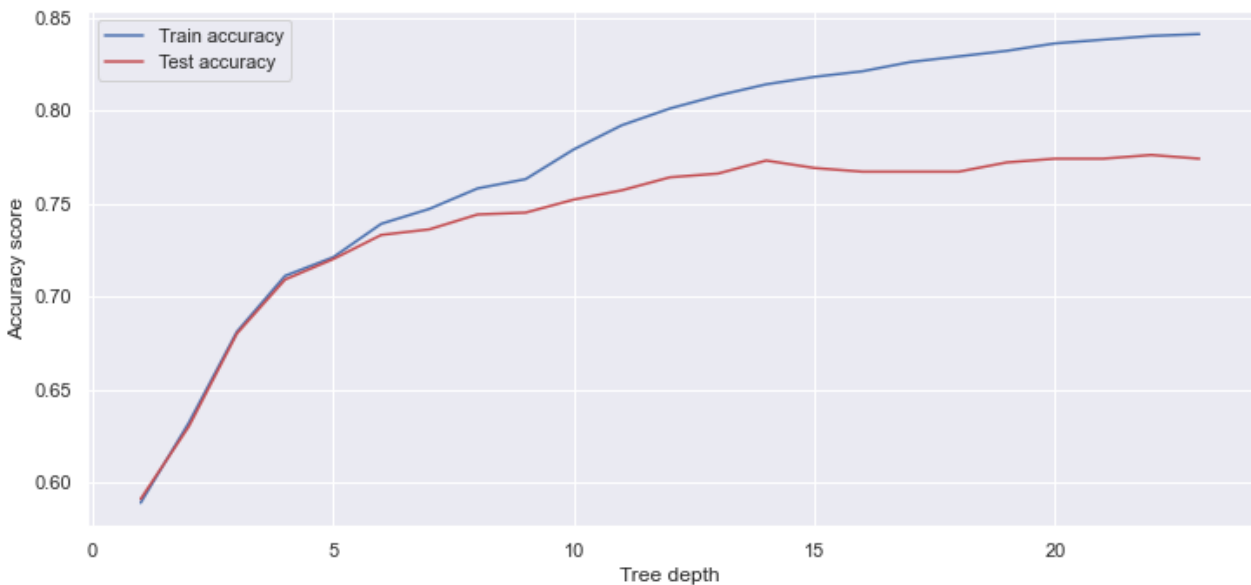
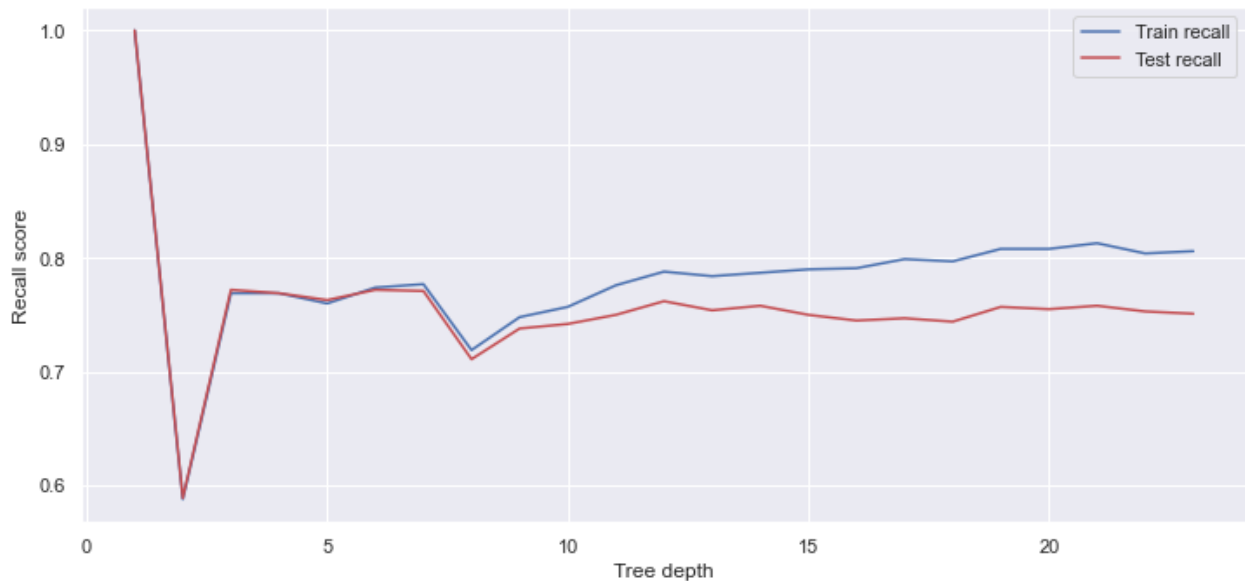
```
In [113]: # identify the optimal tree depth for given data
max_depths = list(range(1, 24))
# create empty list for train results
train_recall_results, train_accuracy_results = [], []
# create empty list for test results
test_recall_results, test_accuracy_results = [], []
# iterate over max depths
for max_depth in max_depths:
    # instantiate a DecisionTreeClassifier
    dt_clf = DecisionTreeClassifier(criterion='entropy', max_depth=max_depth, random_state=42)
    # fit to the training data
    dt_clf.fit(X_train, y_train)
    # find what the model predicts for the training data
    y_train_pred = dt_clf.predict(X_train)
    # calculate relevant statistics for training data
    train_recall, train_accuracy = summary(y_train, y_train_pred)
    # add data to list of train results
    train_recall_results.append(train_recall)
    train_accuracy_results.append(train_accuracy)
    # find what the model predicts for the test data
    y_test_pred = dt_clf.predict(X_test)
    # calculate relevant statistics for test data
    test_recall, test_accuracy = summary(y_test, y_test_pred)
    # add data to list of test results
    test_recall_results.append(test_recall)
    test_accuracy_results.append(test_accuracy)

# set up the plots
fig, (ax1, ax2) = plt.subplots(nrows=2, figsize=(12,12))

# specify the recall plot
ax1.plot(max_depths, train_recall_results, 'b', label='Train recall')
ax1.plot(max_depths, test_recall_results, 'r', label='Test recall')
ax1.set(xlabel='Tree depth', ylabel='Recall score')
ax1.legend()

# specify the accuracy plot
ax2.plot(max_depths, train_accuracy_results, 'b', label='Train accuracy')
ax2.plot(max_depths, test_accuracy_results, 'r', label='Test accuracy')
ax2.set(xlabel='Tree depth', ylabel='Accuracy score')
ax2.legend()

plt.show()
```



In these plots we're looking for the point just before the train accuracy diverges from the test accuracy, indicating that the train accuracy is improving *more* because the model knows the training set *too well* (overfitting).

From the accuracy score, this would appear to coincide with a tree depth of at least 6, but that point on the recall score graph could arguably be anywhere between 8 and 12. We'll compromise with a max tree depth of 10.

```
In [114]: # instantiate a DecisionTreeClassifier
dt_clf = DecisionTreeClassifier(criterion='entropy', max_depth=10, random_state=SEED)
# fit to the training data
dt_clf.fit(X_train, y_train)
# make predictions for the test data
y_train_pred = dt_clf.predict(X_train)
y_test_pred = dt_clf.predict(X_test)
# show relevant statistics
report(y_train, y_train_pred, y_test, y_test_pred)
```

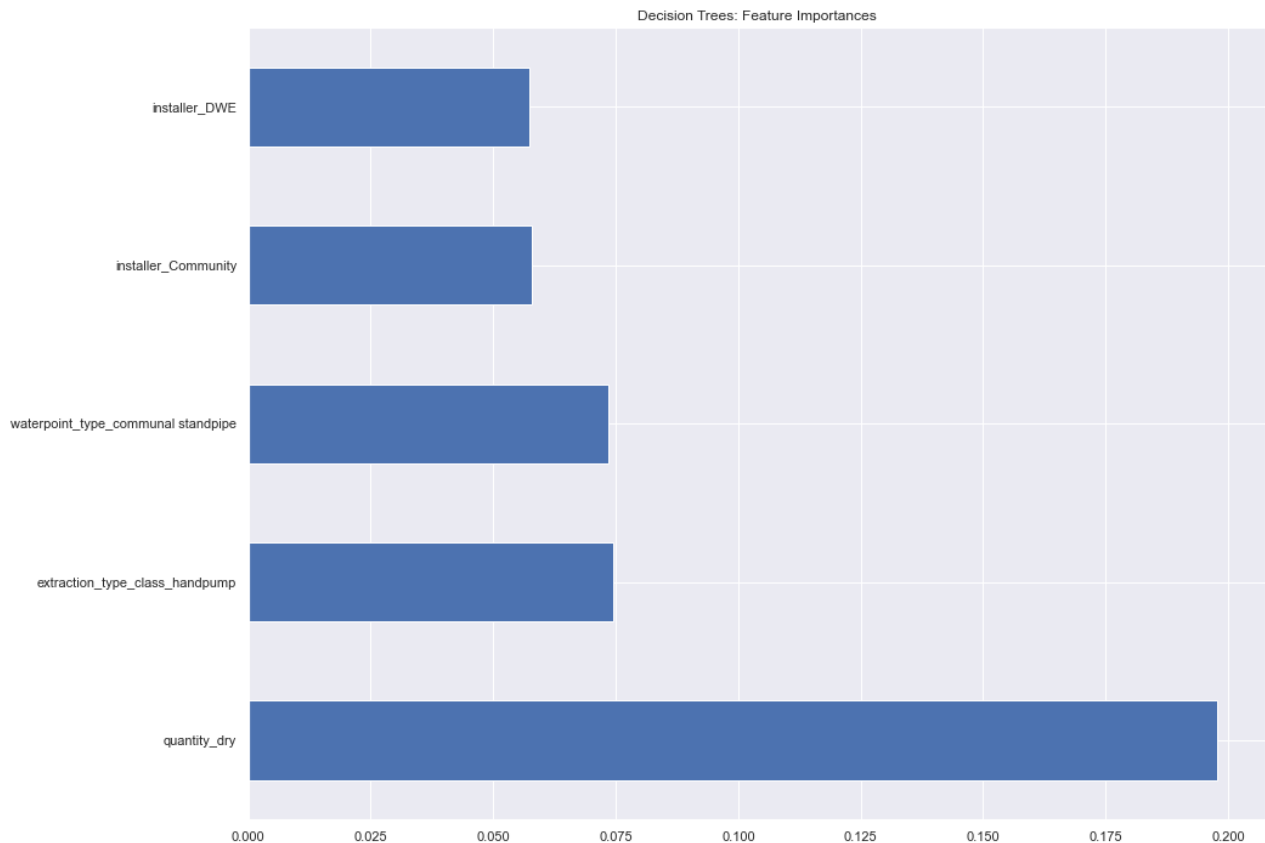
TRAIN SET	recall	accuracy
	0.757	0.779

TEST SET	recall	accuracy
	0.742	0.752

We've sacrificed recall for accuracy, which runs counter to our goals, and this model is not better than the baseline.

Let's use "feature importances" to observe which features mattered most to this model.

```
In [115]: pd.Series(dt_clf.feature_importances_, index=X_train.columns).nlargest(5).plot(kind='barh', title='Decision Trees: Feature Importances');
```



Bagged trees

This model will train on different subsamples of the training data. We'll set the max depth to 8 and start with 20 estimators.

```
In [116]: # instantiate a BaggingClassifier
bt_clf = BaggingClassifier(
    DecisionTreeClassifier(criterion='entropy', max_depth=10), n_estimators=20, random_state=42)
# fit to the training data
bt_clf.fit(X_train, y_train)
# make predictions for the test data
y_train_pred = bt_clf.predict(X_train)
y_test_pred = bt_clf.predict(X_test)
# show relevant statistics
report(y_train, y_train_pred, y_test, y_test_pred)
```

```
TRAIN SET      recall      accuracy
          0.807      0.804
```

```
TEST SET      recall      accuracy
          0.777      0.766
```

This model performs the best so far in either statistic.

Let's perform a similar analysis to what we did with tree depth earlier to see what the optimal number of estimators

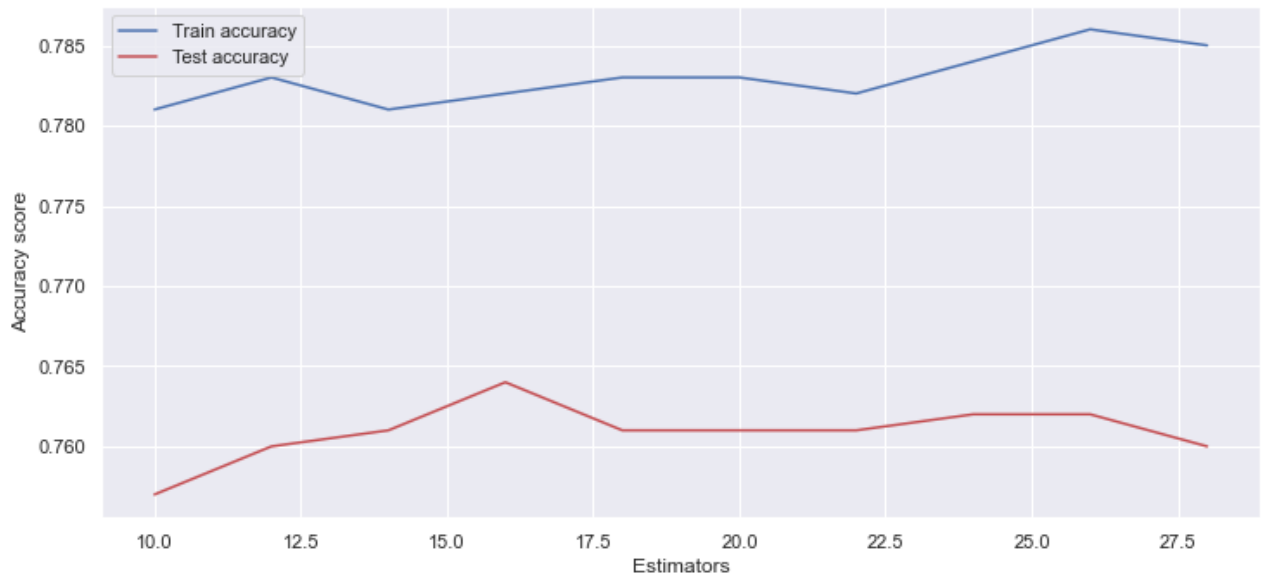
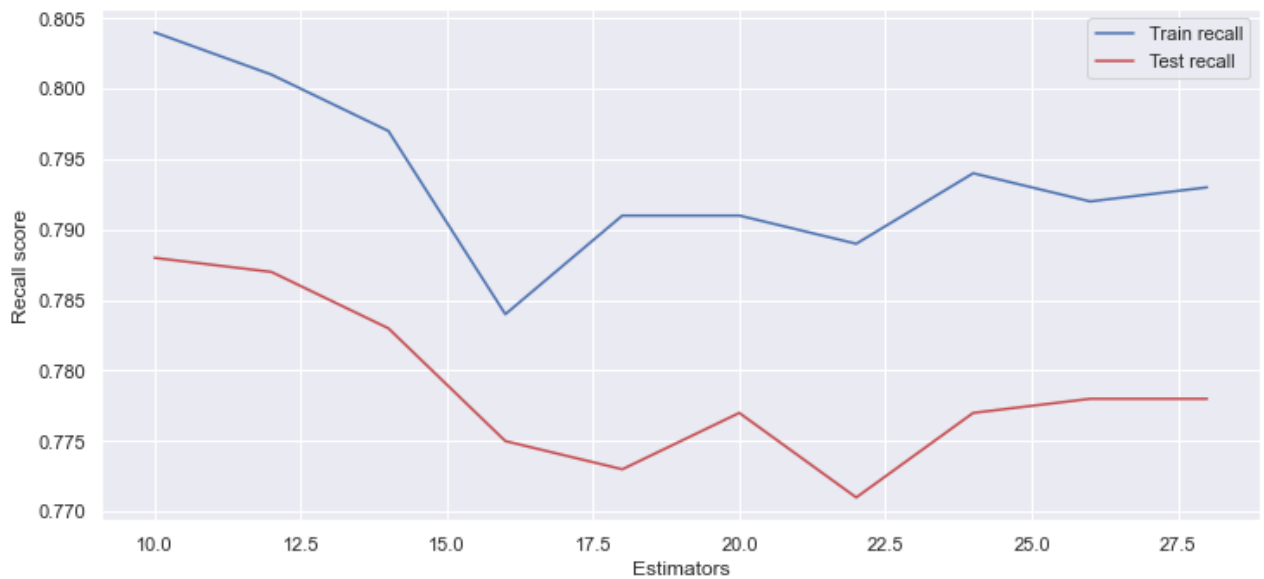
```
In [117]: # identify the optimal tree depth for given data
estimator_list = list(range(10,30,2))
# create empty list for train results
train_recall_results, train_accuracy_results = [], []
# create empty list for test results
test_recall_results, test_accuracy_results = [], []
# iterate over max depths
for estimator in estimator_list:
    # instantiate a BaggingClassifier
    bt_clf = BaggingClassifier(DecisionTreeClassifier(criterion='entropy', max_depth=8),
                               n_estimators=estimator, random_state=SEED)
    # fit to the training data
    bt_clf.fit(X_train, y_train)
    # find what the model predicts for the training data
    y_train_pred = bt_clf.predict(X_train)
    # calculate relevant statistics for training data
    train_recall, train_accuracy = summary(y_train, y_train_pred)
    # add data to list of train results
    train_recall_results.append(train_recall)
    train_accuracy_results.append(train_accuracy)
    # find what the model predicts for the test data
    y_test_pred = bt_clf.predict(X_test)
    # calculate relevant statistics for test data
    test_recall, test_accuracy = summary(y_test, y_test_pred)
    # add data to list of test results
    test_recall_results.append(test_recall)
    test_accuracy_results.append(test_accuracy)

# set up the plots
fig, (ax1, ax2) = plt.subplots(nrows=2, figsize=(12,12))

# specify the recall plot
ax1.plot(estimator_list, train_recall_results, 'b', label='Train recall')
ax1.plot(estimator_list, test_recall_results, 'r', label='Test recall')
ax1.set(xlabel='Estimators', ylabel='Recall score')
ax1.legend()

# specify the accuracy plot
ax2.plot(estimator_list, train_accuracy_results, 'b', label='Train accuracy')
ax2.plot(estimator_list, test_accuracy_results, 'r', label='Test accuracy')
ax2.set(xlabel='Estimators', ylabel='Accuracy score')
ax2.legend()

plt.show()
```



It's not exactly clear what difference tuning this parameter will make. We'll keep the number of estimators at 20.

Random Forest

We'll use those same values for a random forest model.

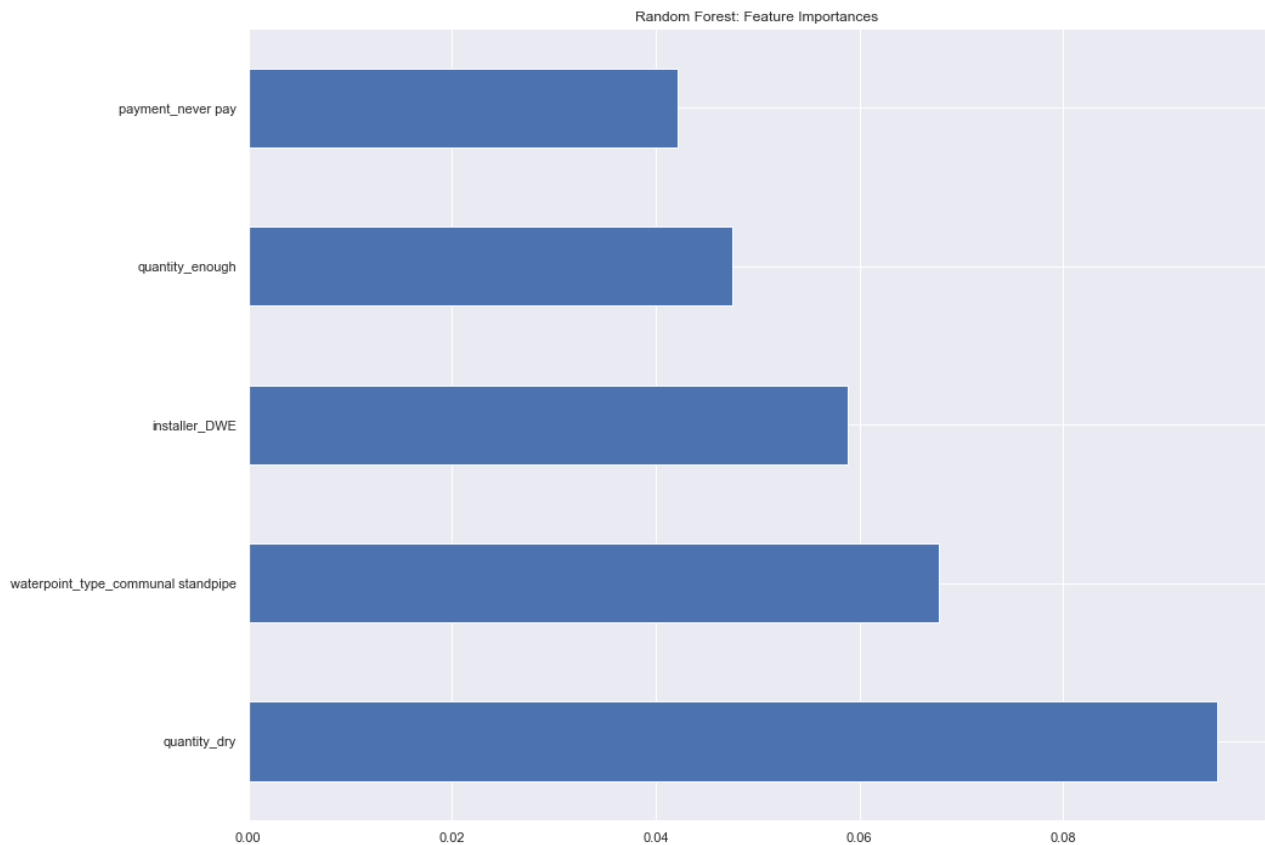
```
In [118]: # instantiate a RandomForestClassifier
rf_clf = RandomForestClassifier(n_estimators=20, max_depth=10, random_state=SEED)
# fit to the training data
rf_clf.fit(X_train, y_train)
# make predictions for the test data
y_train_pred = rf_clf.predict(X_train)
y_test_pred = rf_clf.predict(X_test)
# show relevant statistics
report(y_train, y_train_pred, y_test, y_test_pred)
```

```
TRAIN SET    recall    accuracy
          0.83      0.802
```

```
TEST SET     recall    accuracy
          0.809     0.763
```

This model almost imperceptibly diminishes accuracy but improves recall by about 3%. We'll consider this our new frontrunner. Let's see what features this model considered most important.

```
In [119]: pd.Series(rf_clf.feature_importances_, index=X_train.columns).nlargest(5).plot(
    kind='barh', title='Random Forest: Feature Importances');
```



GridSearchCV

This will in theory take the best of all worlds from a decision tree classifier.

```
In [120]: # set parameters for grid search
dt_param_grid = {
    "criterion": ["gini", "entropy"],
    "max_depth": [None, 6, 8, 11, 15],
    "min_samples_split": [2, 3, 5, 8, 11],
    "min_samples_leaf": [1, 2, 4, 6, 9],
}
# instantiate a GridSearchCV
gscv = GridSearchCV(DecisionTreeClassifier(random_state=SEED), dt_param_grid, cv=3, return_train_score=True)
# fit to the training data
gscv.fit(X_train, y_train)
# make predictions for the test data
y_train_pred = gscv.predict(X_train)
y_test_pred = gscv.predict(X_test)
# show relevant statistics
report(y_train, y_train_pred, y_test, y_test_pred)
```

TRAIN SET	recall	accuracy
	0.797	0.826

TEST SET	recall	accuracy
	0.75	0.773

This is a little better than the best accuracy so far, but it sacrifices too much recall in order to get there.

Adaboost

This model is the classic "adaptive boosting" algorithm, which iteratively prioritizes fixing the previous iterations' mistakes.

```
In [121]: # instantiate an AdaBoostClassifier
ad_clf = AdaBoostClassifier(random_state=SEED)
# fit to the training data
ad_clf.fit(X_train, y_train)
# make predictions for the test data
y_train_pred = ad_clf.predict(X_train)
y_test_pred = ad_clf.predict(X_test)
# show relevant statistics
report(y_train, y_train_pred, y_test, y_test_pred)
```

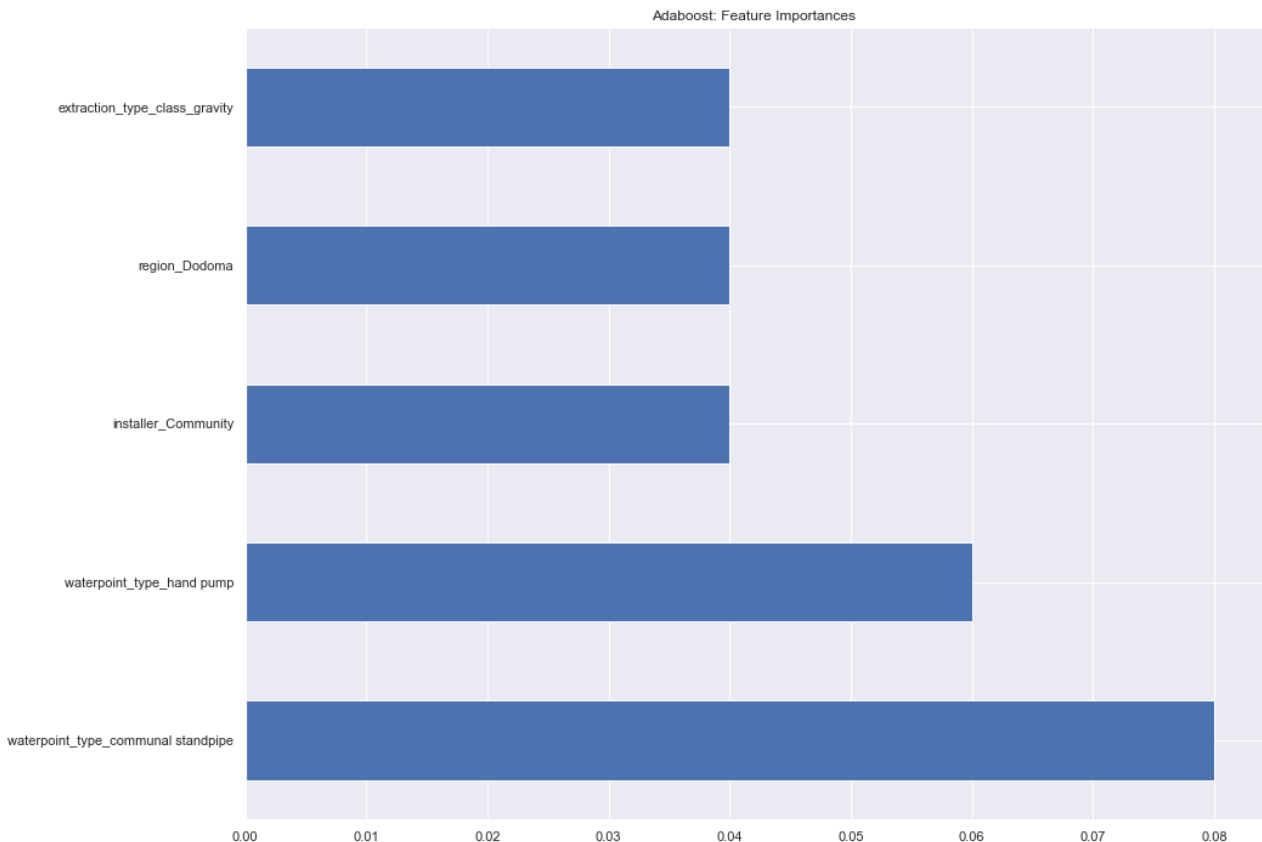
TRAIN SET	recall	accuracy
	0.774	0.744

TEST SET	recall	accuracy
	0.781	0.75

This model is good but falls short of the frontrunner in both statistics.

We'll look at what features this algorithm considered most important.

```
In [122]: pd.Series(ad_clf.feature_importances_, index=x_train.columns).nlargest(5).plot(kind='barh', title='Adaboost: Feature Importances');
```



Gradient Boost

This boosting algorithm iteratively learns in a slightly different way from Adaboost, instead using calculus and gradient descent.

```
In [123]: # instantiate an GradientBoostingClassifier
gb_clf = GradientBoostingClassifier(random_state=SEED)
# fit to the training data
gb_clf.fit(X_train, y_train)
# make predictions for the test data
y_train_pred = gb_clf.predict(X_train)
y_test_pred = gb_clf.predict(X_test)
# show relevant statistics
report(y_train, y_train_pred, y_test, y_test_pred)
```

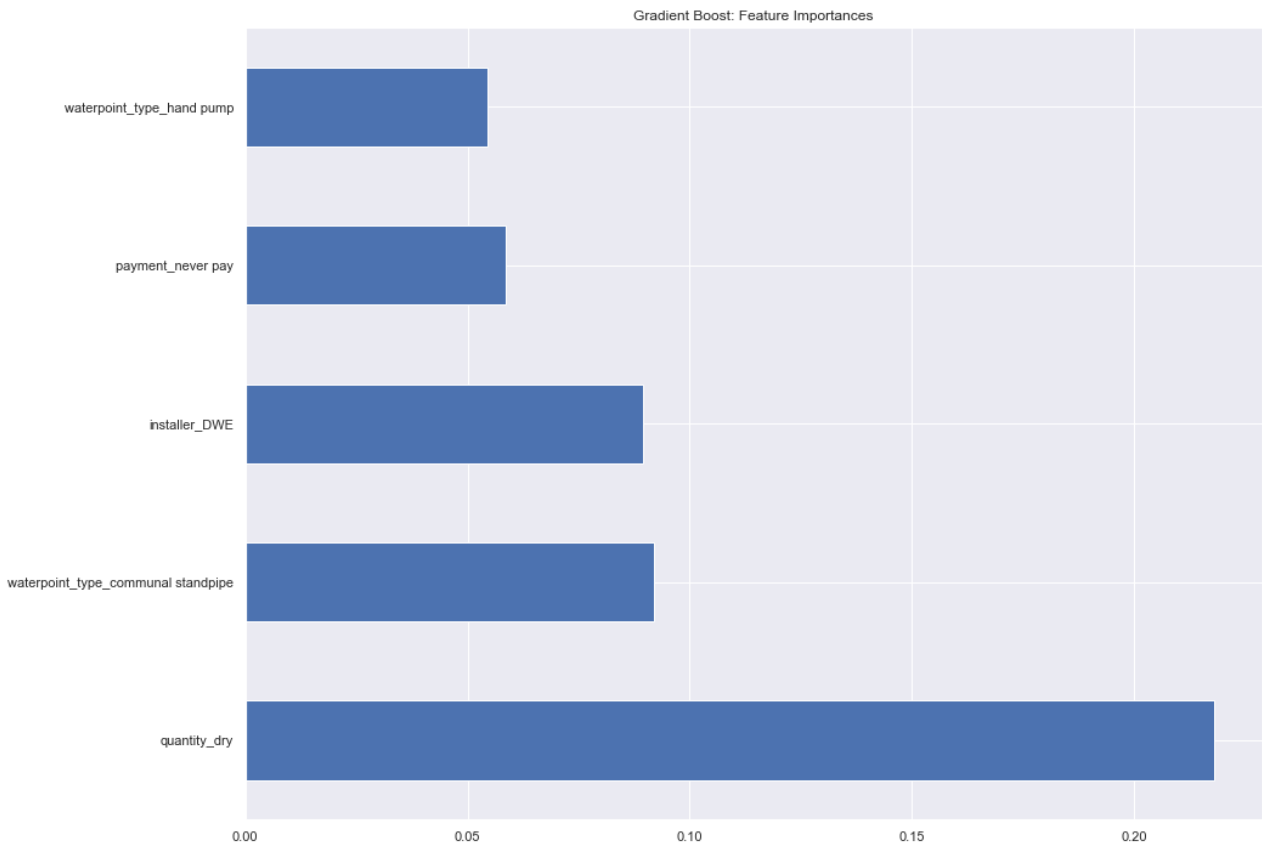
```
TRAIN SET    recall    accuracy
          0.779      0.76
```

```
TEST SET     recall    accuracy
          0.786      0.757
```

This model is just a little better than the previous boost model but also falls short of the frontrunner in both statistics.

Here are the feature importances.

```
In [124]: pd.Series(gb_clf.feature_importances_, index=x_train.columns).nlargest(5).plot(kind='barh', title='Gradient Boost: Feature Importances');
```



XGBoost

XGBoost, or "extreme gradient boost", should produce results as good as or better than any other gradient boost algorithm. Let's see.

```
In [125]: # instantiate an XGBClassifier
xg_clf = XGBClassifier()
# fit to the training data
xg_clf.fit(X_train, y_train)
# make predictions for the test data
y_train_pred = xg_clf.predict(X_train)
y_test_pred = xg_clf.predict(X_test)
# show relevant statistics
report(y_train, y_train_pred, y_test, y_test_pred)
```

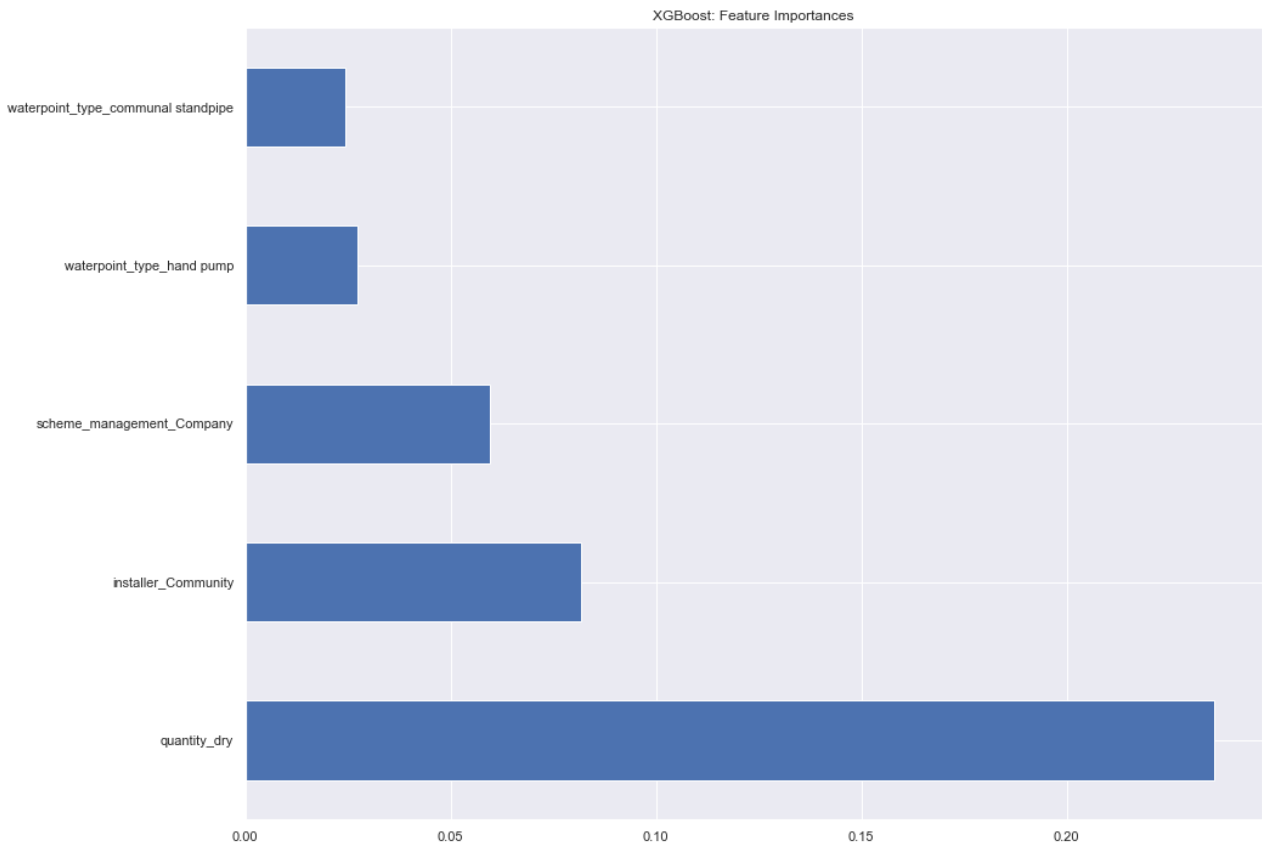
```
TRAIN SET    recall    accuracy
          0.818      0.829
```

```
TEST SET    recall    accuracy
          0.777      0.783
```

This may be the best of the boost models, and it gives the best accuracy score by about 1%, but we do not consider it the best, since it sacrifices too much in the recall score (compared to the current frontrunner, the random forest model).

Let's look at the feature importances.

```
In [126]: pd.Series(xg_clf.feature_importances_, index=x_train.columns).nlargest(5).plot(kind='barh', title='XGBoost: Feature Importances');
```



In the end, the random forest model performed the best according to our goals, which prioritized recall of the non-functional wells over accuracy.

Evaluation

Feature importances

Five of the models provided us with "feature importances". In four of those (including the favorite), *quantity_dry* was identified as the most important feature to the model. There was also a practical consensus among the models that provided feature importances that *waterpoint_type* was the second most important feature and *installer* was the third.

Let's discuss these features in turn.

Water quantity

Let's look at the distribution of water quantity values and compare how functional the wells with those values were.

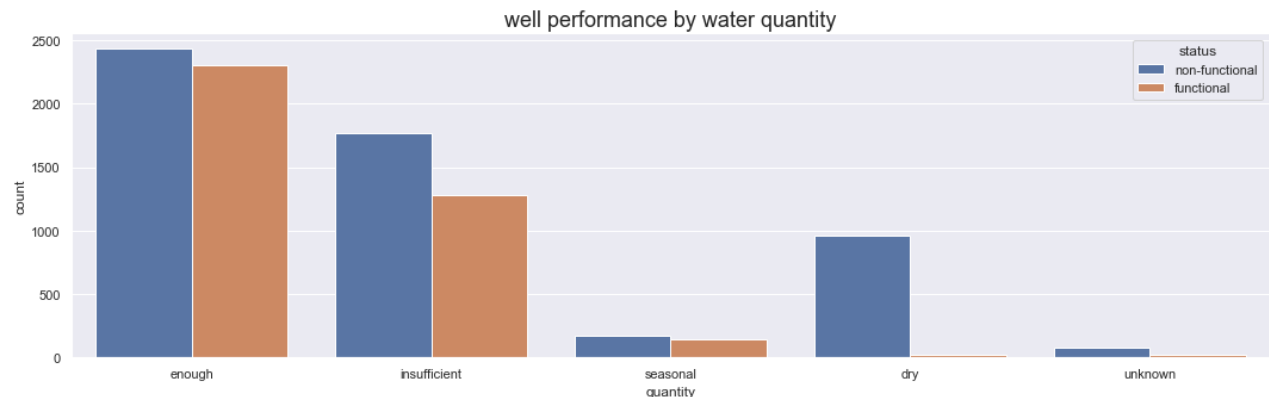
```
In [127]: df.quantity.value_counts()
```

```
Out[127]: enough      4738
insufficient    3050
dry            983
seasonal       316
unknown        95
Name: quantity, dtype: int64
```

```
In [128]: compare_vals('quantity', ['dry', 'unknown', 'insufficient', 'seasonal', 'enough'])
```

```
dry 2%
unknown 19%
insufficient 42%
seasonal 45%
enough 49%
```

```
In [129]: # set figure
fig, ax = plt.subplots(nrows=1, ncols=1, figsize=(18, 5))
# plot well performance
ax = sns.countplot(x='quantity', hue="status", data=df)
# title
ax.set_title('well performance by water quantity', size=18);
```



About 11% of the wells had a "dry" quantity of water, and virtually **all** of those wells are non-functional. The performance of these wells is clearly and distinctly worse than wells of all other water quantity designations, even "insufficient".

Waterpoint type

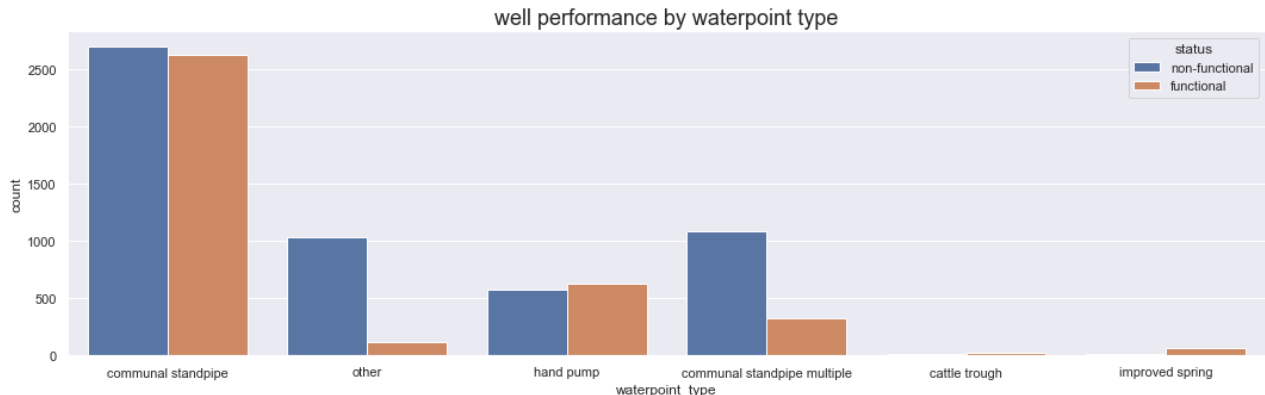
```
In [130]: df.waterpoint_type.value_counts()
```

```
Out[130]: communal standpipe      5312
communal standpipe multiple    1417
hand pump                      1195
other                           1151
improved spring                  77
cattle trough                     30
Name: waterpoint_type, dtype: int64
```

```
In [131]: compare_vals('waterpoint_type', [
    'other', 'communal standpipe multiple', 'communal standpipe', 'hand pump', 'cattle t'])
```

```
other 10%
communal standpipe multiple 23%
communal standpipe 49%
hand pump 52%
cattle trough 67%
improved spring 79%
```

```
In [132]: # set figure
fig, ax = plt.subplots(nrows=1, ncols=1, figsize=(18, 5))
# plot well performance
ax = sns.countplot(x='waterpoint_type', hue="status", data=df)
# title
ax.set_title('well performance by waterpoint type', size=18);
```



There aren't really enough examples of cattle trough and improved spring types to draw hard conclusions, and unfortunately we don't know what "other" even means. We also don't know exactly what the "multiple" designation indicates.

Therefore cautiously draw two conclusions about waterpoint type:

1. "Multiple" communal standpipes are inferior to whatever non-multiple standpipes are.
2. Greater efforts should be made to record the waterpoint type so that the failure of the "other" category can be better understood.

Installer

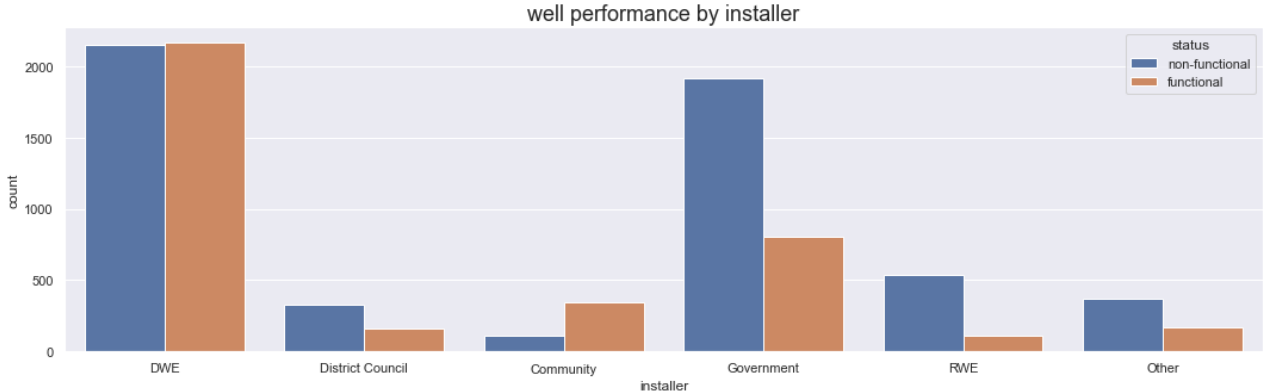
```
In [133]: df.installer.value_counts()
```

```
Out[133]: DWE              4313
Government        2726
RWE                649
Other               544
District Council   489
Community          461
Name: installer, dtype: int64
```

```
In [134]: compare_vals('installer', ['Government', 'Other', 'District Council', 'DWE', 'Community'])
```

```
Government 30%
Other 32%
District Council 33%
DWE 50%
Community 75%
```

```
In [135]: # set figure
fig, ax = plt.subplots(nrows=1, ncols=1, figsize=(18, 5))
# plot well performance
ax = sns.countplot(x='installer', hue="status", data=df)
# title
ax.set_title('well performance by installer', size=18);
```



A cursory search did not reveal just what RWE or DWE are. All we can say is that, insofar as an installer option exists, the data here should guide that choice.

Recommendations

In all three of the important features we discussed, we don't know exactly how much control the government has in choosing among the available options. Insofar as the government has a choice, they should ...

1. place wells in areas with sufficient water quantity
2. favor handpump and communal standpipe waterpoint types over communal standpipe *multiple* waterpoint types
3. have wells installed by the community

However (and wherever) the wells are placed, the government should commit more resources to monitoring wells that ...

1. are placed in areas of low water quantity
2. are of communal standpipe *multiple* waterpoint type
3. were installed by an organization other than the community or DWE (especially wells installed by RWE)
4. were installed before 1985

In all instances, the government should also monitor wells that ...

5. have some or many unknown data points, particularly in the areas of water quantity, waterpoint type, and installer.

Further inquiry

Much of this data was unusable that may have otherwise been rather helpful. We would encourage better data gathering especially in the areas relevant to our recommendations — water quantity level, waterpoint type, and installer identity.

In some cases, it appeared that the "other" or "unknown" labels for a column corresponded with well failure. From this trend we have surmised that perhaps there is some other explanatory variable that correlates highly with well failure and lack of information. This is difficult to demonstrate in modeling, however. More reliable data could help with this.

It would also be helpful to know more about what constraints exist on the choices we've presented. If, for example, we knew how much it cost to implement the different types of wells, we could do additional analysis on which wells were more effective *per dollar spent*. Or it could be helpful to know how the government works with various installers,