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Improving Primary Key Detection with Machine Learning

Bachelor Thesis

submitted by

JANEK PRANGE

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First Examiner : Prof. Dr. Ziawasch Abedjan

Second Examiner : Prof. Dr. Sören Auer

Supervisor : Prof. Dr. Ziawasch Abedjan



DECLARATION

I hereby affirm that I have completed this work without the help of third parties and only with the sources and aids indicated. All passages that were taken from the sources, either verbatim or in terms of content, have been marked as such. This work has not yet been submitted to any examination authority in the same or a similar form.

Hannover, 03.07.2022	
	 Ianek Prange

The discovery of primary key candidates and unique columns in general is an important problem. While primary keys are required to efficiently interact with data stored in a table, various information can be obtained from the uniqueness of a column. The current algorithms to find unique columns work in a linear runtime and are therefore not fast enough for large tables.

In this thesis, I propose a new method to detect unique columns. It uses a machine learning model to predict which columns are probably unique. Only positive guesses will be checked for duplicates using a conventional algorithm. With this method, the time it takes to detect all unique columns is reduced by more than $65\,\%$ for large tables with more than $10\,000\,000$ rows.

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ACRONYMS

AI Artificial Intelligence

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MOTIVATION

Tables are a way to store and organize data that continues to grow in importance. Be it SQL databases, No-SQL databases or simple Excel tables, the ability to organize data and present it concisely is at the core of many projects.

In the best case, the structure of the dataset has already been considered before creating it in order to define which data type can occur in a column, whether values can be left empty and which columns act as primary keys.

However, there are cases where this information is not available for various reasons; either because the data had to be saved quickly and there was no capacity for reasonable planning, or because the data is no longer available in the original format.

In this case, it is an important task to recover the missing metadata as accurately as possible. While information such as the data type and the existence of empty values is comparatively easy to find within a linear runtime, identifying primary keys is more difficult.

One challenge is to distinguish between primary key candidates, which are characterized by the fact that they do not contain duplicates, and practically usable primary keys. An example for the is a column containing comments from users. Even though this column may not contain duplicates, text that is on average 100 characters long is not practically suitable as a key.

Another problem is the time it takes conventional algorithms to determine if a column contains duplicates. The at best linear runtime of these algorithms becomes a problem as the tables become very large and contain several million rows.

This thesis will try to improve the solution to the second problem.

2.1 METRICS

During the experiments in Chapter 5, the performance is measured with different metrics.

Table 2.1: Definition of true and false negatives and positives

		Prediction			
	Positive				
Actual	Negative	True Negative	False Positive		
	Positive	False Negative	True Positive		

Accuracy is calculated by dividing the number of correct guesses by the total number of guesses.

Precision is defined as the number of true positive guesses divided by the total number of positive guesses. It is a good measurement when false positives are associated with a high cost.

Recall is defined as the number of positive guesses divided by the number of actual positives. It is a good measurement when false negatives are associated with a high cost

F1 is calculated by dividing the product of precision and recall by their sum. It is an important measurement when both precision and recall are important and the number of positives and negatives is very unbalanced, for example if there are 99 % negatives.

2.2 MACHINE LEARNING

Machine learning is a part of the field of Artificial Intelligence (AI). The focus is on extracting information from large amounts of data, with algorithms gradually improving themselves to mimic human learning [2].

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The field of machine learning is itself divided into two main parts, Classical learning and Deep learning[7].

2.2.1 Deep Learning and Neural Networks

Deep Learning includes various techniques that use neural networks. In the simplest form, neural networks consist of different so-called nodes, which are arranged as layers[9].

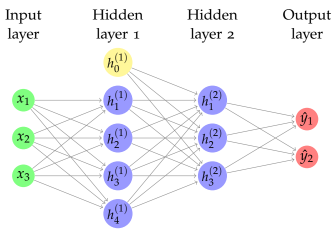
The first layer, also known as the "input layer", is the layer the input vector is applied to. If the input exceeds a certain value, the neuron is activated, i.e. it passes on an output.

In the next layers, which are called "hidden layers", the outputs from all the nodes of the previous layer and a "bias layer" are computed in each node and the output is passed on. The bias layer consists of a single value that influences all nodes on a layer.

Finally, the last layer outputs the result of the computation. How strong each previous node influences the value of the current node is what the neural network is trained on.

Neural networks are particularly good at recognizing patterns in large unordered data, such as images, videos, and audio tracks. Examples of this include facial recognition or verifying hand signatures[5].

Figure 2.1: A simple neural network with three input nodes and two output nodes.



2.2.2 Classical Learning

Algorithms in the field of classical machine learning are largely based on statistical or probabilistic methods, which originated as pattern recognition in the 1960s. It has the advantage of being a fast method that does not require much computing power compared to deep learning algorithms[6].

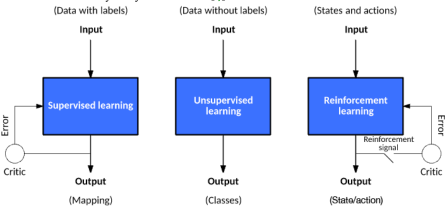
2.2.3 Categories of Machine Learning

Both deep learning as well as classical learning are dived into different categories which are based on the kind of task and trainingsdata the machine learning models have to work with.

Unsupervised learning is used on unlabeled data. Its main use is to organize the data or to make it more manageable. Tasks in this domain include clustering, which categorizes data based on similar data points, or assoziation, which tries to find relationships between variables in a dataset[1].

Supervised learning on the other hand makes use of labeled data. This comes with the disadvantage that the structure of the data as well as the desired results have to be known[6]. Even though it increases the effort of the preparation of the trainingsdata, it also raises the accuracy of the model since it is trained to produce exactly the desired result.

Figure 2.2: This picture shows the different types of machine learning and the way they are trained[4].



2.3 NAIVE ALGORITHMS FOR FINDING PRIMARY KEY CANDI-DATES

There are two basic ways of finding primary key candidates. Both of them are traversing each column and comparing its values to detect duplicates.

The first method works by traversing a column, forming the hash value of each row and saving this hash value in a suitable data structure. If the hash value of a row is already present, the algorithm aborts because it is clear that the column is not unique. This algorithm runs in a time of O(n) as every value in the column has to be checked once to conclude that it is a unique column.

The other method operates by sorting the column in a first step and then comparing each row with its neighbors. If two rows are the same, the column is not unique. This algorithm requires a runtime of $O(n * \log(n))$ for the sorting operation and a runtime of O(n) to check every value of the column.

2.4 USED PACKAGES AND LIBRARIES

The code in this thesis was implemented with Python 3.10.2. To interact with the data, the library pandas was used.

To train the machine learning models, the library Auto-Sklearn which is build on top of Sci-Kit Learn was used.

PROBLEM STATEMENT

Primary key candidates are columns which do not have any duplicate values and can therefore be used to identify a row in the table. For the purposes of this thesis, a column that contains a None/Null value is considered non-unique as it can not be used as a primary key.

The runtime of a conventional naive algorithm is in O(n) or $O(\log(n))$ as explained in Section 2.3. This becomes a problem when the tables have several million rows.

The naive algorithm that is used for the experiments in Chapter 5 is a sorting based method with the specialty of aborting if one of the values is None/Null since the column can not be used as a primary key.

The reason the hashing based method was not used for the experiments was to be able to reliably carry out the efficiency experiments. Although the sorting based method is less efficient, it enables the experiments to be set up in a way that enables the machine learning model to make the correct prediction without decreasing the runtime of the naive algorithm.

PROPOSED METHOD

4.1 OVERVIEW

In this thesis I present a method to increase the efficiency of finding unique columns in a table. The method is based on a machine learning model which uses the first few rows of each column to guess if it will have any duplicate values. Each positive guess will subsequently be validated using a conventional naive method.

The proposed method works in three steps. First, the features are extracted from the first rows of the table. After that, the model tries to predict the existence of duplicate values from the features. Finally, the columns which are unique according to the model are checked with a naive method.

4.2 EXTRACTING FEATURES

The feature extraction is necessary as explained in Section 2.2 to get a table like Table 4.1...

The code in Listing 4.1 prepares each column to produce such a feature table. The feature extraction of the proposed method works in different steps which are executed one after the other.

The variable column contains the first n rows of the column in the table where n is the input size of the method.

First all columns which contain duplicate values in the first rows are sorted out in line 1 and 2 by setting the feature "Duplicates" to 1 and all other features to 0.

In line 5 to 12 the remaining columns are checked in order to determine whether they are sorted or not. During this step, it is possible that an error occurs because two values in the column can not be

Table 4.1: An example for a feature table which is used by the model to predict unique columns. The following values are possible in the column "Data Type": 0: The inspected rows contain a duplicate value. 1: The column contains only integer. 2: The column contains only text. 3: The column contains only boolean values (which does rarely occur without containing duplicates). 4: The column contains a mix of different types.

Duplicates	Data Type	Sorted	Min. value	Max. value	Mean	Std. Deviation	Avg. string length	Min. string length	Max. string length
1	0	0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
0	1	1	7.0	562.0	290.706	187.489	0.0	0.0	0.0
0	1	0	1.0	62.0	28.941	21.496	0.0	0.0	0.0
0	1	0	611.0	946.0	789.118	107.904	0.0	0.0	0.0
0	1	1	1.0	17.0	9.0	5.050	0.0	0.0	0.0
0	2	0	0.0	0.0	0.0	0.0	86.25	78.0	92.0
0	2	1	0.0	0.0	0.0	0.0	78.75	60.0	96.0
0	2	0	0.0	0.0	0.0	0.0	123.032	51.0	296.0
0	2	1	0.0	0.0	0.0	0.0	94.130	46.0	204.0
0	3	0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
0	4	0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

compared. This is mostly the case if there is an empty/None value in the column.

In the following lines, a distinction is made between the different types of values.

If the column contains only boolean values, the feature "Data Type" is set to 3 while all other features stay on 0 in line 13 to 16. Although it is very unlikely that a column contains exclusively boolean values without any duplicate values, there are tables in the gittables dataset to which this applies.

Line 17 to 23 handle columns that contain exclusively numeric values. The "Data Type" feature is set to 1 in line 18 and the four features specific to numeric values are extracted in line 19 and 20. The three remaining features are set to 0 in line 22.

Finally, the columns that contain exclusively string or a mix of different types are handled in the remaining lines.

First, the features for numeric values are set to o in line 25. After that, the average of the length of every item in the column is formed

together with the minimum and maximum length. At the end the "Data Type" feature is set to 2 in line 39.

If there is any value in the column which is not a string, a ValueError is raised in line 32, which leads to the "Data Type" feature being set to 4 in line 42 and the string specific features set to 0.

4.3 TRAINING THE MODEL

How was the model trained? What settings where used for the training and why?

Listing 4.1: This code shows how the features are extracted from a column. This process is repeated for each column; the result forms the feature table which is the input for the machine learning model. The variable column contains the first *n* rows of the column where *n* is the input size of the model.

```
1 if has_duplicates(column):
     return [1, 0, 0, 0, 0, 0, 0, 0, 0, 0]
   result = [0, 0, 0]
3
   # check if entries are sorted
4
   try:
5
     if all(column[i+1] >= column[i] for i in range(len(column)-1)):
6
         result[2] = 1
7
8
     if all(column[i+1] <= column[i] for i in range(len(column)-1)):</pre>
         result[2] = 1
9
   except TypeError:
10
     # mostly this means the column contains None/NaN values
11
12
   if only_bool(column):
13
     result[1] = 3
14
     result += [0, 0, 0, 0, 0, 0, 0]
15
     return result
16
  if only_numeric(column):
17
18
     result[1] = 1
     result += [min_value(column), max_value(column),
19
                 mean_value(column), std_deviation(column)]
20
     # values for strings
21
     result += [0, 0, 0]
     return result
23
   # values for numbers
   result += [0, 0, 0, 0]
25
26
       length_list = []
27
       for value in column:
28
           if isinstance(value, str):
29
                length_list.append(len(value))
30
           else:
31
                raise ValueError("Not a String")
32
       if len(length_list) == 0:
33
           average = 0
34
       else:
35
           average = sum(length_list)/len(length_list)
36
       minimum = min(length_list)
37
       maximum = max(length_list)
38
       result[1] = 2
39
       result += [average, minimum, maximum]
40
41
   except ValueError:
       result[1] = 4 # mixed column, mostly None/NaN value
42
       result += [0, 0, 0]
43
   return result
```

EXPERIMENTS

5.1 EXPERIMENT SETUP

The following experiments where conducted on a server of the university. The machine uses 514 GiB working memory and an AMD EPYC 7702P 64-Core processor. A graphic card was not necessary for the experiments.

5.2 CORRECTNESS

The correctness of the model is probably the most important metric to determine its usability. While a false positive is not a major problem for the correctness because each positive guess is verified (see Chapter 4), a false negative leads to a primary key candidate being ignored.

In this section, different experiments will be conducted to determine which parameters have an influence on the correctness of the prediction. Additionally, in Section 5.2.5 the columns which led to false guesses by the model will be examined.

5.2.1 Experiment Data

The experiments where performed on the GitTables dataset, which is a large corpus of relational tables extracted from CSV files in GitHub[3]. However, not the whole dataset was used but only a subset of tables split into a training and a test dataset.

The training dataset is used to train every model which was tested in the following experiments including the efficiency experiments. It is a subset of the GitTables dataset with 10 000 tables which where selected by traversing the GitTables dataset and skipping tables which are too small.

Every table in it has to have at least 100 rows and 3 columns to ensure the feature extraction described in Section 4.1 does not become the naive algorithm with extra steps. This could be a problem as searching for duplicates with the naive algorithm in the first n rows of each column is part of the feature extraction.

The test dataset was generated the same way as the training dataset. By traversing the GitTables and skipping every table which is too small or part of the training dataset a collection of 5000 tables with 57 211 columns and an average of 184 rows was generated. It was used for every experiment apart from the one in Section 5.2.2 where a dataset of 30 000 tables with 307 030 columns and an average of 277 rows was used.

5.2.2 Comparing models with different input sizes

As described in Section 4.2, the proposed method extracts features from the first rows of a table and uses a machine learning model to predict if a column has any duplicate values based on those features.

In this experiment, I compare different models which use the first 5, 10, 20 and 50 rows of each column to extract features. A model with an input size larger than 50 rows was not feasible as the tables used for training and testing had a minimum size of 100 rows. With a bigger input, the feature extraction would end up performing like the naive algorithm as checking the first n rows for duplicates with the naive algorithm is part of the feature extraction.

Each model was trained for 5 hours on the training dataset. During the experiment, the test dataset with 5000 tables that was described in Section 5.2.1 was used to test each model.

Table 5.1 shows the results of this experiment, which demonstrates that the input size has a large influence on the quality of the prediction. While none of the models has any false negative guesses, the number of false positive guesses decreases with an increasing input size.

This experiment shows that an increase in the input size of the model does have a great impact on the number of false positive guesses. However, since each positive guess by the model is verified using the naive algorithm, the number of false guesses has no effect on the quality of the final prediction of the proposed method. What is however effected by it is the efficiency of the method; this is explored further in Section 5.3.4.

Another important finding of this experiment is that even with a small input size of only 5 rows, the model has not made any false negative guesses. As the negative guesses of the model are not checked, it is important for them to be correct to ensure the overall correctness of the proposed method.

Table 5.1: The result of the correctness experiment comparing models with different input sizes. Each of the models was trained for 5 hours on the training dataset described in Section 5.2.1. The experiment was conducted on the test dataset with 30 000 tables.

Mod	lel Input Size	Accuracy	Precision	Recall	F1
	5	0.704	0.269	1.0	0.424
	10	0.833	0.394	1.0	0.566
	20	0.910	0.547	1.0	0.707
	50	0.970	0.783	1.0	0.878

5.2.3 Altering the training time

The Auto-Sklearn library, which was used to train the machine learning models, automatically searches for the best learning algorithm and optimizes it as described in Section 2.4. Since this process takes time to run, in theory the performance of the model increases with higher training time.

In this experiment, different models with an input size of 10 rows are being trained for different amounts of time on the training dataset. Subsequently, the experiment is conducted for each model on the test dataset with 5000 tables.

The Table 5.2 presents the results of this experiment. With a training time of one minute the performance of the machine learning model is indeed slightly worse as there are some false negative guesses.

However, apart from the false negative guesses, the models with a training time of one and two minutes have a slightly better performance than the other models.

This experiment very clearly demonstrates that the machine learning model does not need a long training time to find unique columns. Furthermore, it shows that there might be advantageous to train a model multiple times and compare them to find the best performing, since a longer training time does not translate to higher performance.

Table 5.2: The result of the correctness experiment comparing models which were trained for different amounts of time on the training dataset. The experiment was conducted on the test dataset with 5000 tables.

Training Time [min]	Accuracy	Precision	Recall	F1
1	0.847	0.369	0.996	0.538
2	0.847	0.369	1.0	0.540
5	0.823	0.337	1.0	0.504
10	0.823	0.337	1.0	0.504
20	0.823	0.337	1.0	0.504
40	0.823	0.337	1.0	0.504
60	0.823	0.337	1.0	0.504
120	0.823	0.337	1.0	0.504
180	0.823	0.337	1.0	0.504
300	0.823	0.337	1.0	0.504

5.2.4 Summary

The experiments in this section demonstrate that the predictions made by the model have a sufficiently high accuracy. Models which have been trained for more than one minute had a recall of 1 and therefore no false negative guesses. This is a very good result as false negative guesses would lead to unique columns which are ignored.

Furthermore, the models in the experiments produce less than twice as much false positive guesses as true positive guesses and even less for models with a larger input size. This is important as false positive guesses reduce the efficiency, which is explored further in Section 5.3.

5.2.5 Examine columns which led to false guesses

The greatest weakness of the proposed method are false guesses. False positive guesses lead to a reduced efficiency because more columns need to be checked with the naive algorithm. False negative guesses on the other hand result in primary key candidates being ignored which reduces the correctness. It is therefore important to examine the columns which lead to false guesses to improve the model if possible.

False positive guesses occur very often as the model is primarily trained to avoid any false negative guesses. The experiment in Section 5.2.2 has shown that depending on the input size between 27% and 78% of the positive guesses are false positives.

The false guesses are unfortunately mostly unavoidable as they are caused mainly by empty cells which are located after the input rows of the model in otherwise unique columns. As the column would be a primary key candidate without these missing values, there is no possible change which would improve the correctness of the model in this case.

Another example for a column leading to a false positive guess is one containing the name of authors. Since the model only sees short strings without duplicates in these columns, there is no good way for the current implementation to recognize the column as non-unique. However, it would be possible to additionally include the column heading as a feature to enable the model to learn that the column with the names of authors is more likely to contain duplicate values.

False negative guesses occurred only in one of all the correctness experiments, namely with the model which was trained for only one minute in Section 5.2.3. And even then the false negatives were rare, and the corresponding columns contained only negative numbers or numbers smaller than 1. While these columns are negative, they are not suited very well as primary keys.

5.3 EFFICIENCY

Next to the correctness, the efficiency of the proposed method is the most important metric to determine its feasibility. The main question is if or from what table size the proposed method is faster than the naive method. This becomes even more interesting as each positive guess of the model has to be verified using the naive algorithm to increase the accuracy.

The following experiments explore the efficiency of the proposed method in comparison to the naive algorithm and which parameters have the greatest influence on it.

5.3.1 Experiment Data

The experiments in this section were conducted on a set of generated tables to control the size of the table as well as the number of unique and non-unique columns. A small example of such a table can be seen in Table 5.3.

The generated tables each have 10 columns and between 100 and 100 000 000 columns. To ensure the correct prediction by the model, the columns where generated in a specific way. The unique columns are evenly incrementing for the first 50 rows, while the first two rows of the non-unique columns contain the same value. The rest of each column contains distinct incrementing values which are mixed up to increase the time the sorting based naive algorithm takes to find unique columns.

Table 5.3: A table generated for the efficiency experiment. The columns o and 1 do not contain any duplicates, the columns 2, 3 and 4 do. To guarantee that the model guesses the unique and non-unique columns correctly, the unique columns are evenly incrementing for the first 50 rows, while the duplicate value of the non-unique columns is in the first two rows.

Index	Column 0	Column 1	Column 2	Column 3	Column 4
0	0	0	100	100	100
1	1	1	100	100	100
2	2	2	93	93	93
3	3	3	45	45	45
:	:	:	:	:	÷
48	48	48	89	89	89
49	49	49	39	39	39
50	91	91	60	60	60
51	77	77	49	49	49

5.3.2 Base experiment

The first experiment explores the efficiency of the proposed method compared to the naive algorithm. The generated tables that were used contained 3 unique and 7 non-unique columns.

Figure 5.1 and Table 5.4 show that for tables with up to 100 000 rows, the naive algorithm takes only a fraction of a second and is therefore

faster than the proposed machine learning model. However, since the model takes a roughly constant time of half a second to compute its prediction, it becomes faster as the table size surpasses one million rows.

The column "Model: Validation" in Table 5.4 additionally illustrates that the validation time of the proposed method is proportional to the number of positive guesses by the model. This highlights the importance of as few false positive guesses as possible, because each false positive guess unnecessarily increases the runtime through the required validation and therefore decreases the efficiency.

In conclusion, this experiment illustrates that for large tables loading the dataset and checking the columns for duplicates with the naive algorithm takes the most time. Possibilities to reduce the loading time will be explored in Section 5.3.3. While a more efficient naive algorithm is not part of this thesis, Section 5.2.2 and 5.2.5 deal with the question of how to decrease the number of false positive guesses.

Figure 5.1: This plot shows the total runtime of the naive algorithm and the proposed method to find the unique columns. The tables were saved in a CSV format.

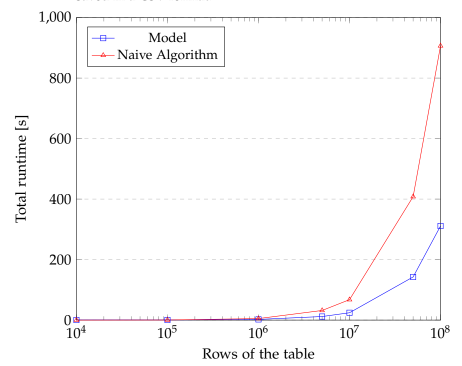


Table 5.4: The result of the efficiency experiment conducted on a table generated with 3 unique and 7 non-unique columns saved as a CSV file.

The experiment was conducted on a model with an input size of 10 rows which was trained on the training dataset.

Rows	Model: Loading	Model: Computing	Model: Validation	Model: Total	Naive: Loading	Naive: Computing	Naive: Total
100	0.001	1.082	0.000	1.084	0.004	0.000	0.004
1000	0.002	0.449	0.001	0.451	0.002	0.002	0.004
10 000	0.006	0.446	0.007	0.459	0.005	0.022	0.026
100 000	0.045	0.452	0.076	0.574	0.045	0.257	0.302
1 000 000	0.434	0.454	1.369	2.260	0.427	4.623	5.050
5 000 000	2.476	0.451	8.824	11.771	2.456	29.341	31.797
10 000 000	4.655	0.446	19.116	24.258	4.606	62.935	67.541
50 000 000	27.298	0.458	114.694	142.650	27.075	381.059	408.133
100 000 000	52.429	0.444	257.904	311.173	52.230	854.417	906.646

5.3.3 *Reduce loading times*

While CSV files are very easy to use, they are not meant to efficiently store large quantities of data. A file format which is substantially more suitable to handle large datasets is the parquet format[8].

It achieves this through the use of various features such as column wise compression, which tends to be more efficient since the values in the same column are usually very similar. This has the additional benefit of enabling the algorithm to only read the required columns which may decrease I/O as only positive guesses need to be loaded for the validation.

Another advantageous property of this format is the concept of row groups, which ensure that a batch of rows is being saved together and can therefore be read together too. This makes it possible to read just the first row group and use these rows as an input for the model.

The Table 5.5 shows the result of the base experiment from Section 5.3.2 repeated with tables generated as parquet files. While the computing time for the model and the naive algorithm remain roughly equal compared to Table 5.4, the loading time is decreased significantly for large tables.

Table 5.6 presents the result for the experiment using the advantages of the file format by loading only the necessary rows and columns. This leads to two loading times for the model. The first time only the first

Table 5.5: The result of the efficiency experiment conducted on a table generated with 3 unique and 7 non-unique columns saved as a parquet file. The experiment was conducted on a model with an input size of 10 rows which was trained on the training dataset.

Rows	Model: Loading	Model: Computing	Model: Validation	Model: Total	Naive: Loading	Naive: Computing	Naive: Total
100	0.004	0.454	0.000	0.458	0.006	0.001	0.006
1000	0.003	0.445	0.001	0.448	0.003	0.002	0.005
10 000	0.004	0.446	0.007	0.457	0.004	0.021	0.025
100 000	0.010	0.447	0.077	0.534	0.010	0.246	0.256
1 000 000	0.040	0.462	1.396	1.902	0.047	4.618	4.665
5 000 000	0.197	1.264	8.751	10.233	0.184	29.033	29.216
10000000	0.482	0.480	18.934	19.938	0.529	62.746	63.275
50 000 000	2.216	0.987	113.886	117.294	2.205	379.468	381.673
100 000 000	4.473	1.549	257.471	263.898	4.395	857.437	861.833

row group is being loaded while the second time only the columns which are unique according to the model are loaded. However, this does not make any difference except for the largest table and even then the total time is hardly changing.

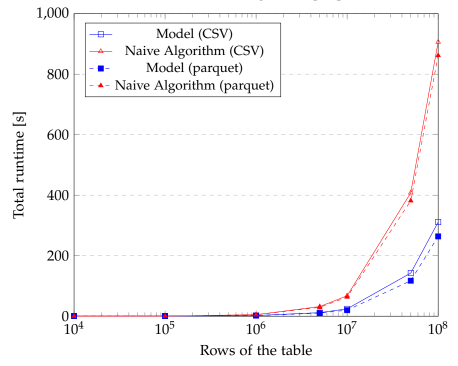
Table 5.6: The result of the efficiency experiment conducted on a table generated with 3 unique and 7 non-unique columns saved as a parquet file. The experiment was conducted on a model with an input size of 10 rows which was trained on the training dataset. During the experiment, only the necessary rows and columns were loaded.

Rows	Model: Loading I	Model: Computing	Model: Loading II	Model: Validation	Model: Total	Naive: Loading	Naive: Computing	Naive: Total
100	0.002	0.458	0.003	0.000	0.463	0.004	0.000	0.004
1000	0.002	0.456	0.003	0.001	0.461	0.003	0.002	0.004
10 000	0.002	0.451	0.003	0.007	0.463	0.004	0.020	0.024
100 000	0.009	1.468	0.006	0.081	1.564	0.009	0.247	0.256
1 000 000	0.032	0.455	0.026	1.356	1.869	0.050	4.666	4.716
5 000 000	0.115	0.463	0.106	8.688	9.371	0.195	29.001	29.196
10 000 000	0.243	0.447	0.258	18.961	19.909	0.544	63.122	63.666
50 000 000	1.183	0.447	1.309	114.895	117.834	2.225	379.731	381.956
100000000	1.602	0.446	2.425	256.993	261.467	4.437	856.737	861.174

In summary, while the reduced loading time does make a notable difference, it is not very large compared to the efficiency gain achieved through the use of the proposed method, as can be seen in Figure 5.2. This could change, however, if the file reading speed would be slower, for example because the data had to be read over the internet. In

this case, reading only the necessary rows and columns and thus decreasing I/O further could make a larger difference too.

Figure 5.2: This plot shows the runtime of the proposed method and the naive algorithm as they search for unique columns in tables of different sizes. I additionally compare the runtime reading the data from CSV files and reading it from parquet files.



5.3.4 Changing the ratio of unique to non-unique columns

The last variable that has an impact on the runtime of the model is the percentage of unique columns in the table. Since every positive prediction by the model has to be verified using the naive algorithm, the total runtime increases the more unique columns the model predicts. The number of unique columns predicted by the model correlates roughly with the number of actual unique columns in the table.

In this experiment, a model with an input size of 10 rows is used on 4 tables, which are saved as parquet files and each have $100\,000\,000$ rows and 10 columns. The difference between these tables is the percentage of unique columns, which range from $60\,\%$ to $90\,\%$.

Table 5.7 shows that nearly every step of the process takes the same amount of time, only the validation step is proportional to the number of unique columns.

In the GitTables dataset, which is used in the correctness experiment, the ratio of unique columns is 10 %. The positive guesses of the model are quite a bit higher since its priority is to avoid false negatives, not false positives. Still, the share of positive guesses during tests on the GitTables dataset is around 30 %, which is low enough to be a clear improvement over the naive algorithm given large enough tables.

Table 5.7: The result of the efficiency experiment where each table has a size of 100 000 000 rows and 10 columns and is read from a parquet file. The only thing that is changing is the number of unique columns.

Unique Columns	Model: Loading	Model: Computing	Model: Validation	Model: Total	Naive: Loading	Naive: Computing	Naive: Total
4	4.489	1.308	344.742	351.127	4.493	861.111	865.603
3	4.473	1.549	257.471	263.898	4.395	857.437	861.833
2	4.445	1.180	171.984	177.881	4.388	862.440	866.828
1	4.568	0.469	87.070	92.244	4.497	856.509	861.006

5.3.5 Summary

The experiments in this section show that the proposed method to find primary key candidates is suitable for some cases. If the tables that will be examined contain mostly viewer than 1000000 rows or the ratio of unique to non-unique columns is too high, the model is probably slower than the naive algorithm. However, on very large tables with 100 000 000 or more rows the model can significantly improve the overall runtime.

Section 5.3.4 additionally demonstrates that for a high efficiency it is important to decrease the number of false positive guesses made by the model as much as possible.

CONCLUSION

6.1 POSSIBLE APPLICATIONS

The proposed method is not suitable as a standalone application the way it is implemented as there are too many cases where the efficiency is far behind the naive algorithm. A more promising possibility would be to use this method as a part of a larger algorithm or program where it would be used only if it has an advantage of the naive algorithm.

Another useful advantage of the proposed method is the fact that it produces a set of possible unique columns in a very short amount of time even for large tables. This would make it possible to very fast display columns which may be primary key candidates and validate the guesses in the background.

It may be possible to transfer the idea behind the proposed method to use a machine learning algorithm which predicts if a column has certain properties (such as uniqueness). This way it could be possible to improve the speed in several areas of data analysis, not just the detection of primary key candidates.

6.2 LIMITATIONS OF THE PROPOSED METHOD

A big limitation of the proposed method is the fact that it only predicts uniqueness for single columns.

Additionally, there is the possibility that a unique column is classified as a non-unique by the proposed method. However, to rule out this possibility, all the columns would have to be inspected with the naive algorithm.

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