# CREATING MASTER-DATA FROM DATASET USING SIMILARITY-SCORES OF TEXT-FIELDS

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## INTRODUCTION

Most conglomerates today have many separate applications and systems (viz. ERP, CRM) where data that crosses organizational departments or divisions can easily become fragmented or duplicated. Reporting critical KPI (Key-Performance-Indicators) for a business accurately becomes difficult. Questions like "How many entities do we govern?", "Which of them are most profitable?", require a cleaned and accurate master dataset. The data-capturing phase itself might lack a standardized approach, resulting in fundamental discrepancies rendering the data unusable for reporting. An incorrect address in the customer-master might mean orders, bills, and marketing literature sent to the wrong address; an incorrect account number in an account master could mean huge fines.

# MOTIVATION FOR THE CASE-STUDY

Generating and maintaining a Master dataset from data collated via various third-party vendors, internal source systems, and data integrated in-cases of acquisitions and mergers at one central data-hub repository is challenging. A 'single version of truth' maintained for each associated entity across the entire organization's data sources, can help orchestrate collaboration between multiple cross-functional channels of the business.

This paper focuses on *masterizing* clinical data in terms of hospitals/sites, that the pharmaceutical client manages. For example- the same site "Kadlec Regional Medical Center", might be reported differently as "Kadlec Clinic Hematology and Oncology" but with the same address, across the client's source systems. Our goal is to identify a *golden entity* (Master Record) to which other duplicate records can be matched, and maintain their *source-to-master linkage* (Cross-Reference). Although industry-standard tools are available (Informatica, Oracle, SAP, etc.) that can be used with third-party collaborators like *Address-Doctor-Service*, or *Dun&Bradstreet* to retrieve the standardized asset data, this case study was intended to prove that open-source code and libraries can produce near-standardized results.



Fig. 1.1. Basic Deduplication example

## LITERATURE SURVEY & IMPLEMENTATION CHOICES

For de-duplicating records and generating a master set, we compute the similarity between two textual strings and determine if they are a probabilistic data match. If two or more records seem to belong to the same *golden entity* i.e. the Master record, they are 'linked' together. The intuition behind this procedure is as follows:

- Within a dataset of n records, we must compare the 1<sup>st</sup> record with the remaining (n 1) records, the 2<sup>nd</sup> record with the remaining (n 2) records, and so on. Thus, there would be  ${}^{n}C_{2} = n \cdot (n 1)/2$  unique combinations to be considered.
- Between 2 different datasets of m and n records each, there would similarly be  $m \cdot n$  unique combinations to be considered.

At an individual combination level i.e. for the participating records, a string-comparison algorithm <sup>[1]</sup> will be used to compute a match-score of the relevant feature-strings. Let *str1*="Kadlec Regional Medical Center" and *str2*="Kadlec Clinic Hematology and Oncology".

1. Edit-distance-based algorithms (ex- Levenshtein) compute the number of character-level operations needed to transform one string to another. More the number of these character addition or subtraction or replacement operations, less is the similarity between the two strings. For example- the Levenshtein distance between str1 and str2 will be 25, and the normalized-similarity will be:

$$\frac{LevenshteinDistance}{MaxLength} = \frac{|Mismatching chars|}{Max(str1.length, str2.length)} \approx 0.325$$

Jaro-Winkler is a similar directional algorithm that checks for characters of *str1* occurring in a window of some size within *str2*.

2. Token-based algorithms (ex- Jaccard-index) will find similar tokens in both string sets. More the number of common tokens (words or n-gram characters), greater is the similarity between the sets.

$$J(str1, str2) = \frac{|str1 \cap str2|}{|str1 \cup str2|}$$

For str1 and str2, using words as tokens the score  $\approx 0.125$ , while using individual character-tokens gives  $\approx 0.558$ .

3. Sequence-based algorithms (ex- Ratcliff-Obershelp similarity) try to find the longest sequences present in both strings. First, remove the longest common substring from both strings, and split the originals into the left and right parts of the common substring. Repeat this recursively for both the left and right parts, until the size of any broken part is less than a default value. The score is twice the number of characters found in common divided by the total number of characters in the two strings.

$$RO(str1, str2) = \frac{2 \cdot |str1 \cap str2|}{str1.length + str2.length} \approx 0.45$$

**4. Cosine-similarity** can be summarized as a widely used NLP technique that uses a matrix of word-embeddings <sup>[2]</sup>: where each cell in a column, represents the weight by which the word associates to that row/attribute. Two words x and y, are first converted to their word-vectors from this word-embedding matrix, and the cosine formula is applied to identify semantic similarity:

$$CosineSimilarity = \frac{x \cdot y}{||x|| \cdot ||y||}$$

Kaitlin Coltin et al. observed that Levenshtein produces results on par with Cosine-similarity, when matching potential duplicate organization names against a master list [4]. Bearing in mind their comparison against cosine-similarity, coupled with the fact that a high volume of our dataset contained junk characters and spelling errors, and considering the anagram-possibility scenario of Jaccard-measure (which could lead to undesired matches), it made sense to use the Levenshtein algorithm. In contrast to their approach, we wanted to identify the unique entities in our dataset with no standard set available. Hence, deduplicating the input by comparing it against itself was vital for our process. Similarly, machine-learning techniques like clustering or classification algorithms, weren't applicable since there isn't a target variable/list to train or test on.

The **RecordLinkage** library in R provides two main functions to generate n\*(n-1)/2 candidates for deduplication within a single dataset (hereafter called the *dedup* function), or m\*n candidates for identifying duplicates between two different datasets (hereafter called the *linkage* function) [5]. The Python equivalent library is limited by the array-size that Pandas can hold when the number of candidate-pairs is ginormous [6]. However, Python's easy-to-use data-wrangling features, ability to invoke a child-subprocess like R-scripts, topped with some deployment-server versioning limitations, led to developing the end-to-end structural pipeline in Python. R is used only for generating match-scores (indirectly by using a pre-compiled C function), since in-memory statistical computations are much faster, and Pandas cannot generate such massive sized-DataFrames in-memory [7].

Since the deployment server supported only Python 2.5x and R 3.4x, we had to refactor the Python code, and reverse-engineer the overall **RecordLinkage** library in R (since it requires R >= 3.5.0). The original core capabilities of the *dedup* and *linkage* functions were maintained, but the cursory code supporting phonetic algorithms, blocking datasets, etc. was removed to speed up the algorithm. The Levenshtein function can be implemented in multiple ways [3], but we picked the source code written in C by Joe Conway, Murat Sariyar, and Andreas Borg [5] since it is already a part of the package. The source code in C was pre-compiled into its binaries, and reloaded into R using the following commands:

- > R CMD SHLIB levenshtein.c
- dyn.load("levenshtein.so")

Binaries generated in Windows have the .dll (Dynamically Linked Libraries) extension and in Unix the .so (Shared Object) extension.

Once the *dyn.load*() function loads the binaries and the symbols within, the Levenshtein function can be invoked from within R, to return the number of characters replaced/added/removed to make the strings match.

## **ARCHITECTURE**

#### a. PREPROCESSING THE RAW-DATA

The incoming name and addresses for clinical sites and hospitals were gathered from more than 5 source systems owned by the global pharmaceutical client and contained more than 30 countries in the data. The ETL process is done in Informatica PowerCenter, and the Site name-address details are loaded into a single table, which is then leveraged to generate a CSV input file for the algorithm's input. The algorithm outputs a Master dataset and the Cross-References dataset, which are then loaded back into the Database.

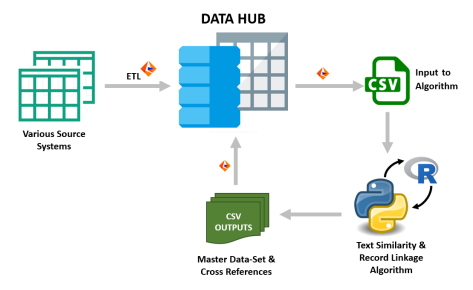


Fig. 2.1.1. High-level ETL (Extract Transform Load) pipeline architecture

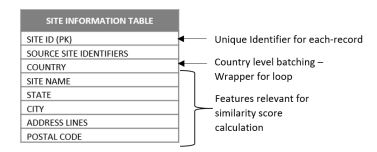


Fig. 2.1.2. Structure of the algorithm's input (Site-data compiled from multiple sources)

For now, the data is filtered on the below conditions before feeding it to the algorithm script:

- Standardized Country-Names is a mandatory field across the dataset and cannot be NULL.
- For a record to be 'linked' to another near-duplicate record, the sum of the match-score of features must be greater than a threshold. Hence, input records must have a minimum number of NOT NULL features to be able to pass the threshold score: *Site-Name*, *Postal-Code*, *State*, *City*, *Address-Line*.

The following section outlines these implementations in a bit more detail:

• Batches are formed using a wrapper for-loop on the Country-Names; firstly because this field is standardized in the preprocessing ETL phase using ISO-standard translation tables, to act as the most

- reliable field amongst all others, and secondly, 2 or more duplicate Site records would implicitly belong to the same country.
- Features relevant to calculating a match-score are used to sort the data into minibatches of size less than or equal to some threshold.
  - This ensures that even though the data might contain junk characters or spelling errors, each minibatch itself can contain a high volume of duplicates already, leading to higher compression.
  - A Row-Number ID is assigned to records, to be used in the cross-reference table for backtracking. (Refer Fig. 2.1.3.)
- The raw CSV is ingested into a pandas DataFrame in UTF-8 encoding to ensure Non-Latin scripts are handled.
  - It is then cleaned using simple ETL functions like removing punctuation marks, replacing NULL with blanks, and ensuring the index of the DataFrame is the integer row-number column generated in the previous step.
  - Address-Fields 1 to 3, are concatenated into a single column, and individually dropped.

Sr. No.	Country	Site-Name	State	City	Addr-Line-1	Addr-Line-2	Addr-Line-3	Zip code
1001	Algeria	Centre Hospitalo Universitaire de Batna	Batna	Batna	Allées Mohamed Boudiaf	NULL	NULL	05000
1002	Algeria	Centre Hospitalier Universitaire Tlemcen	Tlemcen	Tlemcen	Boulevard Mohamed V	NULL	NULL	13000
1003	Algeria	Centre Hospitalo Universitaire de Constantine	Alger	Alger	11 BP, Colonel Amirouche	NULL	NULL	16000
1004	Algeria	Centre Pierre et Marie Curie	Alger	Sidi M'Hamed	Place du 1er Mai 1945	Centre Hospitalier Universitaire Mustapha Pacha	NULL	16000
1005	Algeria	EPH Mascara	Mascara	Mascara	Mascara	NULL	NULL	29000
1006	Argentina	Hospital Universitario Austral	Buenos Aires	Buenos Aires	Avenida Juan D. Peron 1500	NULL	NULL	01629
1007	Argentina	FUNDALEU - Fundacion para combatir la Leucemia	Ciudad Autonoma Buenos Aires	Ciudad Autonoma Buenos Aires	José E.Uriburu 1450	NULL	NULL	1114
		•		•	•	•	•	•
	•	•	•	•	•	٠	•	•
1499	Argentina	Hospital Italiano de Buenos Aires	Ciudad Autónoma de BuenosAires	Ciudad Autonoma de Buenos Aires	Calle Tte Gral Juan Domingo Peron 4190	Department of Oncology	NULL	1199
1500	Argentina	Hospital Britanico de Buenos Aires	Ciudad Autonoma Buenos Aires	Ciudad Autonoma Buenos Aires	Perdriel 74	NULL	NULL	1280
1501	Argentina	Hospital Britanico de Buenos Aires	Ciudad Autonoma Buenos Aires	Ciudad Autonoma Buenos Aires	Perdriel 74	NULL	NULL	1280

Fig. 2.1.3. Sorted dataset splits into minibatches with a high volume of potential duplicates

#### b. RECURSIVE PROCESSING

This algorithmic approach will first pass minibatches of a fixed size into the *dedup* R-function and generate deduplicated master-datasets. These deduplicated master-datasets would be compared against each other using the *linkage* R-function. This is similar to the conventional level-order traversal of a binary tree using a queue, but in reverse, until each record is compared against every other. The motivation here is to prevent overuse of RAM, due to in-memory candidate pair computations.

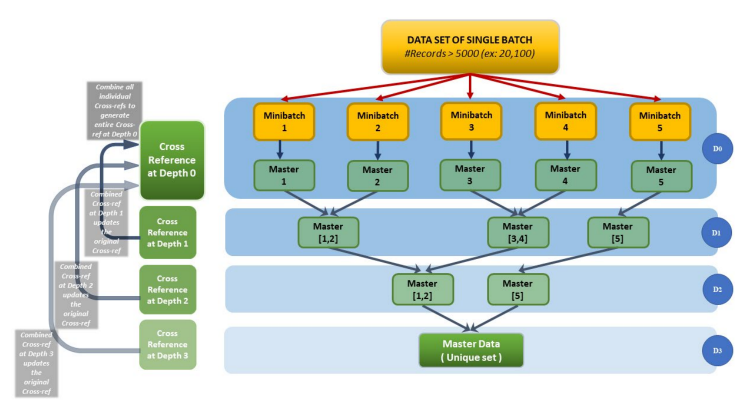


Fig. 2.2.1. Pipeline for the Recursive approach

At Depth=0, the number of minibatches will be:

$$m = \frac{EntireBatchSize}{MAXSIZE} + 1$$

For each of these m iterations, a CSV file will be generated after the *dedup* function, and added to a queue of CSV file names. A cross-reference DataFrame will be maintained for the entire batch that will keep getting updated during each step of the process.

For example- for an incoming batch of 20,100 records:

- i. There will be 5 minibatches considering each of size 5,000.
- ii. Rather than computing match-scores for 201,994,950 candidate pairs\* in a single go, the algorithm *masterizes* 4 minibatches of 5,000 records and 1 minibatch of 100 records using the *dedup* function.
- iii. The pre-sorted data in each minibatch leads to high volume compressions; on average 80% are identified as duplicates of the remaining 20% Master records, (on average we'd identified 1000 unique masters).
- iv. These 20% Master records are written to a CSV file and the file name is added to a queue of CSVs.
- v. After the 5 iterations, the Cross-Reference at Depth 0 has all 20,100 entries (source-to-master linkages).

<sup>\*</sup>  $(n \cdot (n-1)/2) \approx 201$  Million combinations and 5 features to compare for each combination. Huge DataFrame required: (201 Million rows) x (10 feature columns)

For the subsequent Depths =  $1, 2 \dots (m+1)/2$ , we pop 2 file names at a time from the queue and process them using the *linkage* R-function. The output of each pair is written as a new CSV, and the file name is appended to the same queue. If a second CSV is not present in the current queue to compare against the first CSV, simply write the first CSV as the output dataset of the comparison. Similarly, maintain a Cross-Reference of each depth which will be used to update the Cross-Reference generated at Depth 0, with what has newly been observed as source-to-master linkage. Upon comparing a set of masters amongst each other at depth d, update these newly identified masters into the existing Cross-Reference of the entire batch.

The time taken for recursively processing a large batch is significantly lower than the time that would've been theoretically required for one-shot processing. The following observations were taken by considering minibatches of size 5,000, on an AWS EC2 instance m5.4xlarge (64 GB RAM, and 16 vCPUs- each a single thread on a 3.1 GHz Intel Xeon Platinum 8175M processor):

Input batch-size	Candidate-pairs n(n-1)/2	Minibatches (BatchSize/5000)+1	Time required	Comment
735	269,745	1	5 sec	Single minibatch processed
3,500	6,123,250	1	2 min	Single minibatch processed
5,000	12,497,500	1	5 min	Single minibatch processed
22,882	261,781,521	1	N/A	One-shot processing would theoretically require ≈90 min.  However, the child process itself gets killed (RAM usage exceeds limit)
22,882	261,781,521	5	35 min	5 mastered minibatches created, and recursively processed faster than the theoretical expected time

Fig. 2.2.2. Execution stats of different volumes of input batch

### c. INTERPRETING SCORES AND IDENTIFYING MASTERS

For both, the *dedup* and *linkage* R-functions, we use thresholds to convert the normalized Levenshtein-similarity score into a binary-values to indicate if the feature matches for a candidate pair or not. The address-match score is scaled up by a factor since in many cases, the state, city, and postal code were empty/different but showed up in the address. We sum up all comparison outputs to produce a total score of that candidate pair. If the total score is greater than or equal to TOTAL MATCHES THRESHOLD, this candidate pair is considered for further processing.

Features	Match-score Threshold	Scaling Factor	Max possible Score
Site Name	0.85	1	1
State	0.85	1	1
City	0.85	1	1
Concatenated Address	0.75	3	3
Postal Code	0.85	1	1
Total Matches Threshold	4		7

Fig. 2.3.1. Thresholds/parameters for the match-score computations

i. The output of these R-functions can be interpreted as *the raw universe of potential duplicates* for that minibatch; a DataFrame containing the following columns:

[ Source-Record-Id, Master-Record-Id, Site-Name-Comparison-Score, State-Comparison-Score, City-Comparison-Score, Address-Comparison-Score, Postal-Code-Comparison-Score ]

- ii. The source-record can match against multiple master-records with a total match-score  $\geq 4$ . We choose the best match for incoming source-records based on the highest total score for all its potential master-records (a Greedy Approach).
- iii. There are cyclic cases in these score outputs like-

Record B matches against Record A Record C matches against Record B

Ideally, we should transitively maintain:

Record C matches against Record A

These cyclic occurrences may extend to upwards of 10-15 such transitive linkages, so handling them efficiently is crucial.

iv. Finally, from this list of cleaned-normalized-score-features, using basic set-theory we find the unique list of masters. Consider 'SR\_NUM\_1' as the list of incoming Source-Ids, and 'SR\_NUM\_2' as the Master-Ids to which 'SR\_NUM\_1' should be linked based on match-score.

'SR NUM' of the entire minibatch, will be the universe of records.

Union of 'SR NUM 1' & 'SR NUM 2' will be the universe of potential duplicates (UPD).

Stand-alone records in the current minibatch, are those which do not fall in this *universe of potential duplicates* (Non-UPD).

The final Master-records will be the union of Master-Ids and the Stand-alone Ids identified above.

```
U = \{SR\_NUM\}
SourceIDs = \{SR\_NUM\_1\}
MasterIDs = \{SR\_NUM\_2\}
UPD\ IDs = \{SourceIDs\} \cup \{MasterIDs\}
NonUPD\ IDs = \{U\} - \{UPD\ IDs\}
MasterList = \{NonUPD\ IDs\} \cup \{MasterIDs\}
```

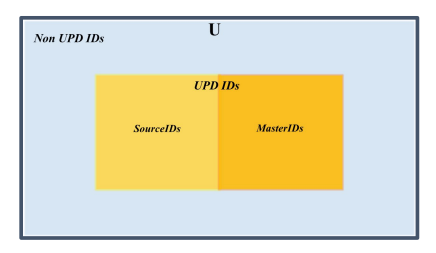


Fig. 2.3.2. Identifying master records by interpreting score output

# **OUTCOMES**

The *masterization* process for a batch finally produces the Master file and a Cross-reference file which contains a translation of each record against either: a parent Master-record, or itself (in case of the record itself being a Master-record). The following table concisely summarizes the functionality of the existing algorithm based on a few scenarios covered for testing:

Country	Test case description	Observation	Status
USA	Batch has size>mini-batch size.  Data should be processed in recursive minibatches	Data processed in recursive minibatches	Pass
Turkey	Batch has special characters in UTF-8 encoding.  Master data should be generated by considering special chars.	Comparison can handle special chars	Pass
UK and Northern Ireland	NULL valued features should match with each other with score=0	Comparisons against NULL values are given a score of 0	Pass
France	Batch with potential duplicates. Master data should look unique.	There are still some potential duplicates due to data-scenarios (refer to Future Scope in the next section)	Fail
Paraguay	Batch has a single record.  The single record should be marked as master itself	The single record is marked as master itself	Pass

Fig. 3.1. Test cases and scenarios covered

The following table gives the compression statistics for a few batches monitored. Compression percent of Raw-data to Master-data is just the percent of duplicates found.

$$Compression~(\%) = \frac{\mid Duplicates \mid}{\mid Input \mid} * 100 = \frac{\mid Input - Unique \mid}{\mid Input \mid} * 100$$

Country	Input Dataset size	Unique Masters identified	Data Compression %
USA	22,881	8,818	61.46
Turkey	2,275	764	66.42
UK and Northern Ireland	1,885	631	66.53
France	1,628	639	60.75
·			•
	<u> </u>		•
Paraguay	1	1	0
Overall	50,937	18,695	63.29

Fig. 3.2. Compression stats for input datasets

In the absence of a reference Master-list of site information that the client governs, it wasn't possible to compute the accuracy/precision of the output. However, the next section will dig into the current shortcomings, which were fairly noticeable.

## **FUTURE SCOPE**

- 1. Lemmatization is the grouping together of a word's different inflected forms to a single item, i.e. it links words having a similar meaning, to one word [8]. Lemmatizing each word in the features, during the preprocessing step itself, can improve match-score computations. Python has a library called NLTK (Natural Language Processing Tool Kit) for this. However, in the case of spelling errors (a high volume in our dataset), the output word would be the same as input since no root word would be found and the overall performance *might* not change.
- 2. Implement an incremental approach to match: an incoming dataset of delta records vs the already identified Master-records present in the Database. Records with Country=NULL should also be handled in this case.
- 3. The scoring output (*raw universe of potential duplicates* within a minibatch) generated by computing each candidate-pair's match-scores can have cyclic occurrences. For now, the following is our strategy, but it has scope for improvement:
  - i. If 1548 matches with 1543 with a total score of 7 (max possible score)
  - ii. But 1543 itself matches with 1541 probably with a lesser score of 4 (lower score)
  - iii. If site-names are different, remove this candidate-pair [1543 vs 1541] from this universe of potential duplicates, thus making 1543 and 1541 as 2 separate master records.
  - iv. If site-names are the same for this candidate-pair [1543 vs 1541], 1541 would be the final parent record for both 1548 and 1543.
- 4. Fine-tune the process of master-record selection; instead of the Greedy Approach explained in the "Interpreting Scores" section (i.e. choosing the very first record amongst the universe of potential duplicates within a country), scan through this universe and check for max-occurrences of features within that subset. The best candidate for the golden Master-record, would be the one having the highest overall combined-similarity score. For ex, for a subset of interlinked candidate-masters: 1501, 1502, 1503, and 1504, consider that 1501 matches against 1502 with a total score of 4, against 1503 with a total score of 5, against 1504 with a total score of 7 and so on:

Candidates	1501	1502	1503	1504	Total Preference
1501	-	4	5	7	16
1502	4	-	6	7	17
1503	5	6	_	4	15
1504	7	7	4	-	18

Fig. 4.1. Better identification of master amongst a set of duplicates

1504 could be the best candidate here for the golden Master-record since it has the highest overall combined-similarity score.

- 5. Address-Fields 1, 2, and 3 can have human errors, ex: addr2 of first record might be same as addr3 of second record, or addr3 may not be present for one record, but could be a huge string for second record, leading to address-comparison mismatch. Improved approach could be to compare each combination of addresses for each address fields, viz. [a.addr1 vs b.addr1], [a.addr1 vs b.addr2], [a.addr1 vs b.addr3], [a.addr2 vs b.addr1], [a.addr2 vs b.addr2], and so on.
- 6. When the state and city are abbreviations rather than full-words, ETL can standardize them by maintaining an ISO-standard translation table for abbreviation vs full-name.

7. Develop and maintain a front-end application to enable business users to look at merge scenarios and take actions themselves: Merge or unmerge 2 different records.

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