Efficient Record Linkage Algorithms Using Complete Linkage Clustering

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Abstract

Data from different agencies share data of the same individuals. Linking these datasets to identify all the records belonging to the same individuals is a crucial and challenging problem, especially given the large volumes of data. A large number of available algorithms for record linkage are prone to either time inefficiency or low-accuracy in finding matches and non-matches among the records. In this paper we propose efficient as well as reliable sequential and parallel algorithms for the record linkage problem employing hierarchical clustering methods. We employ complete linkage hierarchical clustering algorithms to address this problem. In addition to hierarchical clustering, we also use two other techniques: elimination of duplicate records and blocking. Our algorithms use sorting as a subroutine to identify identical copies of records. We have tested our algorithms on datasets with millions of synthetic records. Experimental results show that our algorithms achieve nearly 100% accuracy. Parallel implementations achieve almost linear speedups. Time complexities of these algorithms do not exceed those of previous best-known algorithms. Our proposed algorithms outperform previous best-known algorithms in terms of accuracy consuming reasonable run times.

Introduction

Health agencies keep track of patients’ health information and at the same time records of a patient reside in multiple data sources. All the records of a patient may be needed to accurately diagnose a disease or prescribe medicine for a disease for the patient [1, 2]. Disease evolution, drug discovery and side effects of a drug may require analysis of health records across these data sources [3, 4]. Record linkage, for example, can be used to merge records across educational databases, employment history, and family evolution to analyze an individual’s characteristics. It has also applications in similarity detection in digital documents [5, 6], master data management [7], social networking [8], historical research [9], gene expression [10–13], information science [14], health psychology [15], data mining [16, 17], etc.

Record linkage [18] integrates records across multiple data sources as well as identifies records pertaining to same individuals. Now-a-days millions of records are stored and maintained...
in data sources electronically. Connections among these records provide better understanding of relationships of these data sources. Exact same records exist in multiple databases. Sometimes records get polluted unintentionally due to typing error, similarity in pronunciation, etc. All of these issues make the record linkage problem very challenging and critical. Efficient algorithms are inevitable to address this problem.

Fortunately, a large number of algorithms are available in the real world [19]. A naïve algorithm compares each pair of records to find matches. It may produce expected results but has a high time complexity. Therefore algorithms have been devised to provide best possible results within a manageable time. We have previously proposed single linkage hierarchical clustering based solutions [20] for this record linkage problem. These algorithms provide very fast solutions in finding clusters of individuals with a high accuracy.

In this paper we propose a complete linkage hierarchical clustering based solution for this problem. Single linkage solution works fine for real life applications. But it has a chaining problem. We discuss the problem elaborately in this paper. Our newly devised algorithms not only solve this chaining problem but also assure expected output. We also develop an efficient parallel version of this algorithm. Our experimental results substantiate our claim.

Background and Significance

Record linkage [21, 22] identifies record matches across different data sets even if they have no universal identifier. The problem is to group similar records so that each group contains all records of one individual only. This problem is no more than trivial if the records do not get contaminated. Often errors are introduced unintentionally while typing, due to sound similarity, etc. Every group of similar records can be thought of as a cluster. Every cluster should contain only the records of a single person and it should contain all the records of this person. Several types of clustering algorithms such as $k$-means clustering, fuzzy clustering, hierarchical clustering, graph-based clustering, etc. are widely available [23]. Our proposed algorithms are based on hierarchical clustering [24]. This requires linkage criteria that define how distances are measured between any two clusters. Single linkage and complete linkage clustering are popular in use. In single linkage, the distance between two clusters $A$ and $B$ is computed as the minimum distance between a point (i.e., a record) in $A$ and a point in $B$. In complete linkage, the distance between two clusters $A$ and $B$ is computed as the maximum distance between a point in $A$ and a point in $B$. Therefore single linkage clustering can be thought of as the nearest neighbor clustering and complete linkage clustering can be thought of as the farthest neighbor clustering. In addition to defining the distance between two clusters, we also have to define the distance between two records. There are many distance measures that can be used for the records. Edit distance or Levenshtein distance calculates the number of insertions, deletions and substitutions required to transform one string to the other. (We can think of every record as a string of characters). Manhattan distance computes only the number of mismatches. There exist some other distance calculation methods such as Euclidean distance, maximum distance, etc. We have used complete linkage hierarchical clustering for our algorithms. These algorithms generally use edit distance, reversal edit distance and truncation edit distance calculation methods although our algorithms can support any distance measure. Reversal edit distance and truncation edit distance also use edit distance calculation methods.

Related Works

A naïve or brute force algorithm compares every pair of records and hence takes too much time. There exist a large number of efficient algorithms [25, 26], [27, 28] define data cleansing and record linkage. They also present a literature survey for many proposed or developed
methodologies for entity resolution and record linkage. A relational clustering algorithm uses both attribute and relational information to integrate entities [29]. Discussions about deduplication quality and data linkage measurement involve different linkage processes and issues [30]. Limitations in record linkage algorithms have also been discussed in the literature [31]. The EMH algorithm (based on expectation maximization) provides better decision rules employing probability estimates [32]. There exist some other probabilistic methods for record linkage problems [33, 34]. A hybrid Markov chain Monte Carlo algorithm calculates transitive linkage probabilities across records and uses this information for post-processing procedures such as logistic regression [35]. Relational probability model can solve the citation-matching problem [36]. Records across multiple data sets may contain variations as well as errors [37]. Edit distance calculation has been used widely to compute variations between records [38]. Case patient algorithm includes 'Jaro—Winkler', 'Soundex' and 'weight matching' for distance computation [39]. Record linkage has also applications in record matching [40], text correction [41], substring matching [42], etc. Relational dependencies among different fields improve record linkage processes by reducing errors [43, 44]. Conditional models for record linkage problem can handle varieties of features of input data sets independent of their dependencies [45, 46].

Blocking and indexing have been used extensively for faster computation by removing many unnecessary pair comparisons [47–49]. Traditional blocking, sorted neighborhood indexing, Q-gram-based indexing, suffix array-based indexing, canopy clustering, and string-map-based indexing are popular blocking techniques for reducing comparison space. [50] proposes Q-gram fingerprinting as a blocking technique. It transforms records into bit vectors and filters pairs of bit vectors using multibit trees. FEBRL [51, 52], FRIL [53, 54], Intelliclean [55] are well-known and widely used record linkage algorithms and tools. FEBRL uses three different indexing methods namely standard blocking method, sorted neighbourhood approach, and n-grams. It has a parallel implementation using MPI with python. FRIL is another good tool for record linkage with many options. It employs nested loop join (NLJ) and the sorted neighborhood method as search methods. Hierarchical clustering based solution has been popular for record linkage [56–59]. Given the exponential growth in data sizes, parallel solutions are inevitable [57–63]. Recently developed single linkage hierarchical clustering algorithms outperform these algorithms [20]. In this paper we propose sequential and parallel record linkage algorithms that use complete linkage clustering. These algorithms offer improved accuracies and have the potential of having a greater impact on real world applications.

Methods

We propose sequential and parallel record linkage algorithms, which use complete linkage hierarchical clustering. These algorithms employ single linkage algorithms [20] as a preprocessing step to generate intermediate clusters. Complete linkage method is applied within each of these clusters. We employ some post processing steps to fine-tune the clusters thus generated.

Sequential Algorithm

RLA-CL (Record Linkage Algorithm—Complete Linkage) works in several phases and each of these phases consists of possibly multiple steps. Steps involved in RLA-CL are blackshown in Fig 1.

RLA-CL first sorts the records and identifies duplications. As different data sets may have different numbers and types of attributes, it takes pairs of data sets in which one of them has a subset attribute types of the other. Then the algorithm sorts them using efficient radix sort on
Start

Take a set of data sets and a threshold value

Combine data sets and sort lexicographically

Find exact clusters by merging duplicate records

Generate blocks of records sharing common $k$-mers

Construct a graph by considering records as vertices and connections as edges. Connect a pair of records if the edit distance is $\leq$ the threshold value

Find connected components in the graph

Split each connected component into one or multiple groups such that any pair of vertices within a group has a distance $\leq$ threshold value

Expand each group by bringing back duplicates of each record

Write expanded groups into disk files

Stop

Fig 1. A flow chart describing all steps involved in RLA-CL.

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common attributes. Exact matches will be adjacent in the sorted array. We do this sorting for each pair of data sets meeting our required criteria. We accumulate all of them and eliminate duplicates by merging them into the same clusters. This single phase removes many records from further consideration and shrinks the data sets. A simple example may simplify the working process of this phase. Let $A$, $B$, and $C$ be three input data sets. $A$ has $a$, $b$, $c$, and $d$ as attribute fields, $B$ has $a$ and $d$ and $C$ has $a$, $d$, and $e$. Note that the attributes in $B$ form a subset of the attributes in $A$. Also, the attributes in $B$ form a subset of the attributes in $C$. We sort $A$ and $B$ together; we also sort $B$ and $C$ together. The attributes in $A$ form neither a subset nor a superset of the attributes in $C$. Thus we do not sort $A$ and $C$ together. After sorting the records of $A$ and $B$ data sets, we accumulate duplicate records into clusters. We do the same process for $B$ and $C$ data sets. Then the algorithm merges these two arrays of clusters obtained from merging $A$ with $B$ and $B$ with $C$ data sets. This exact matching phase identifies all possible duplicates and unifies them into clusters. Therefore the remaining phases of the algorithm have to handle these reduced data sets only, which form a subset of the initial data sets. We have shown this phase in Algorithm 1.

**Algorithm 1** Find Exact Clusters

**Input:** A list of records

**Output:** A set of clusters of identical records

1: procedure FINDEXACTCLUSTER
2: for each pair data set { $X$, $Y$ } do
3: if attributes of $X$ is a subset of attributes of $Y$ then
4: Combine records from $X$ and $Y$;
5: Sort lexicographically using radix sort;
6: Merge duplicate records by creating clusters of identical records;
7: Remove duplicate records;
8: end if
9: end for
10: Merge clusters generated from all pairs;
11: return the set of exact matched clusters.
12: end procedure

Exact matching results in clusters of records. From each cluster we pick only one representative for further processing. In this way we make our algorithms independent of the number of input data sets and they can identify similar records within a data set as well as across different data sets.

Comparison between every pair of records is time consuming and impractical. Blocking helps to reduce the number of pairs to be compared. We employ $k$-mers or $k$-substrings of an attribute for blocking. If the blocking attribute contains only English letters, numbers, or alphanumeric values, then we consider only $26^k$, $10^k$, or $36^k$ blocks, respectively. Each block has only those records having at least one $k$-mer of the blocking field in common. If $l$ is the length of the attribute of a record, then this record goes to $(l - k + 1)$ blocks. If two records belong to the same person and if an attribute slightly differs in these two records, then there is a good chance that the two attribute instances will still have a common $k$-mer and hence the two records will fall into at least one block together. We measure distances among records in each block. We generally employ edit distance, reversal distance and truncation distance calculation methods although every suitable distance calculation works perfectly with our algorithm. Edit distance calculates the minimum number of insertions, deletions and substitutions of characters needed to change one string to the desired one. If $S_1$ = “algorithmss” and $S_2$ = “algorithms”, then we can convert $S_1$ to $S_2$ using the following operations: insert ‘o’ at index 3 of $S_1$, replace ‘l’ to ‘t’ at index 5 and delete ‘s’ from index 9 of $S_1$. This algorithm discards many calculations by checking when the distance surpasses the user-defined threshold.
value. Therefore we have to choose a suitable threshold value dependent on our input data accuracy. Threshold value defines the maximum number of errors allowed in the input records. For the above example if the threshold value is not less than 3, then the algorithm integrates them into a single cluster. These steps have been shown in Algorithm 2.

**Algorithm 2** Perform Single Linkage Clustering  
**Input:** A set of exact matched clusters and a threshold value  
**Output:** A set of single linkage clusters  

1: procedure **COMPUTESINGLELINKAGECLUSTER**  
2: Take a record from every exact matched cluster as a representative;  
3: In the next steps by a record we mean a representative record;  
4: for each attribute in a user defined attribute list do  
5: Create blocks of records sharing the same \(k\)-mer;  
6: for each block do  
7: Consider a graph where records are vertices and connections among them are edges;  
8: Connect two vertices if the distance (edit distance is one of the distance calculation methods) between them is at most the user defined threshold value;  
9: end for  
10: end for  
11: Remove multi-edges and self-loops to make the graph simple;  
12: Find connected components of this graph;  
13: return the set of connected components in the above graph.  
14: end procedure

If records are considered as vertices and distances not above threshold value as edges, then we get an undirected graph. We remove multi-edges between pairs of vertices and self-loops to convert it into a simple graph. We find all the connected components of the graph. These connected components are intermediate clusters generated by single linkage clustering method. This is the third phase of our algorithm.

Our next phases work on only records within each cluster. Every cluster typically contains a small number of records integrated by single linkage clustering. Single linkage clustering often traps in a chaining problem. Let \(A, B, \) and \(C\) be records, where \(A = \) ”sweat, exercise, gymnasium” having status, type, and place as attributes, \(B = \) ”sheat, gymnesium” with status and place as attributes and \(C = \) ”heat” having status as the attribute. Let the threshold value be 1. Therefore \(A\) and \(B\) are in one cluster, and \(B\) and \(C\) are in another cluster, but the distance between \(A\) and \(C\) is 2, which is above the threshold value. According to our first three phases all the three records should be considered in the same cluster. Complete linkage removes this problem. It may merge \(A, B\) in a cluster and \(B, C\) in another cluster or \(A, B\) in a cluster and \(C\) in another cluster, and so on. It never merges \(A, B\) and \(C\) in a single cluster.

The fourth phase starts with considering every record in a cluster as a cluster having only one record. Then the algorithm measures distances among each pair of clusters and populating them in 2-\(d\) matrices. From these distances we generate a vector having minimum distances from every single record cluster. The algorithm finds the minimum of them, and if this minimum distance is not above the threshold value, then it merges these two clusters into one cluster and updates the distance matrix and vector. When we calculate the minimum distance for a cluster, we measure distances of the furthest elements between every pair of clusters having this cluster at one side and take the minimum of them. This process continues till the minimum distance does not surpass the threshold value. We eventually get clusters of records of individuals using complete linkage clustering.
Algorithm 3 RLA-CL (Record Linkage Algorithm using Complete Linkage Clustering)

Input: A set of data sets and a configuration file
Output: A set of complete linkage clusters

procedure RLA-CL

Find exact clusters using Algorithm 1;
Compute single linkage clusters by Algorithm 2;

for each single linkage cluster do

Consider every record of the cluster as a single node cluster;
Generate a $2 \times d$ square matrix where each entry contains the minimum distance between pairs of clusters; Each row of the matrix corresponds to a cluster; Generate a vector of minimum distances for each cluster;

while the matrix has more than 1 row do

Merge clusters if the minimum distance between them is no more than the user defined threshold value;
Update the matrix and vector;
end while

Check whether merging is possible among the generated clusters;
Use a priority list to resolve ambiguity in finding a perfect cluster for each record;
end for

Merge these clusters with records from exact matched clusters;
return these complete linkage clusters.
end procedure

The fourth phase easily eliminates the problem of merging all the records in a single cluster generated by the chaining phenomenon. But which cluster should contain which records is now a challenging task. We employ a post-processing phase to fine-tune the generated complete-linkage clusters. We require a user-defined priority list of attributes to complete this phase. We assign each priority attribute a score. We take one record from one cluster and check in which cluster it matches the best. The error-free matching with higher priority attributes, clusters having the highest number of priority attributes, etc. determine the destination cluster. This process meets the user-expectations astonishingly in real world applications. Algorithm 3 describes every step of the algorithm. We can explain the above algorithm using a simple example. Data set A has 3 records 
{("Cade", "Bale", 05011976), 
{"Cade", "Bolt", 05021986}, and 
{"Thor", "Glenn", 12011990}, and data set B has 2 records namely 
{"Thor", "Glenn", 12011990} and 
{"Cade", "Balt", 05011976}. Both of these data sets have first name, last name and date of birth attributes. Let the blocking field be first name; comparing attributes be the first name and last name; the priority field be date of birth; and the threshold value be 1. RLA-CL first accumulates these five records and sorts. It finds four exact matched clusters. Only one cluster 
{"Thor", "Glenn", 12011990}, 
{"Thor", "Glenn", 12011990}] has two records having the same first name and last name. Then the algorithm creates blocks on the first name for all of the four representative records. After blocking and constructing linkages, we find 2 clusters. One is 
{"Thor", "Glenn", 12011990} and the other is 
{"Cade", "Bale", 05011976}, 
{"Cade", "Bolt", 05021986}, 
{"Cade", "Balt", 05011976}. The post processing phase finds an inconsistency: the "Balt" record may go with the "Bolt" record or the "Bale" record since the edit distance value in both cases is 1 and the threshold value is 1. To break this tie, the priority field date of birth helps us to combine the "Balt" record with the "Bale" record. After expanding exact matched records we get 3 clusters 
{"Thor", "Glenn", 12011990}, 
{"Thor", "Glenn", 12011990}, and 
{"Cade", "Bolt", 05021986}.
### Analysis

We analyze the time complexity by aggregating time complexities of all the steps. Step 1 calls radix sort for at most $D^2/2$ data sets, where $D$ is the number of data sets. If $D = 10$, which is very high for real world applications, the sorting algorithm is called at most 50 times. As radix sort is a linear time algorithm, this step consumes a linear amount of time on the number of records contained in those pairs of data sets. Step 1 reduces the number of records significantly in practical applications. Let the initial number of records be $N$ and this reduced number be $N'$. K-mer blocking is typically done on alphabet, number or alphanumeric values which generates $26^k$, $10^k$ or $36^k$ blocks, respectively. If a record length is $l$, then it should be in $(l - k + 1)$ blocks.

To calculate blocking information of all the records, step 2 takes at most $(l' - k + 1)N'$ time, where $l'$ is the maximum length of any blocking attribute. Step 3 is the most time consuming step as it measures distances between records in every block. Let $b$ be the number of blocks, $b_n$ the average number of records in these blocks and $L$ be the maximum aggregated length of common attributes of records. Then this step takes $O(b_n^2 L \tau)$ time, which can be written as $O(b_n N' L \tau)$ as $bb_n = O(N')$. Step 4 scans through the generated graph and finds connected components. This step takes linear time in the number of records and connections, which is $O(N')$. Steps 6 and 7 work on individual clusters that contain small numbers of records. If the number of these clusters is $C$ and each cluster may contain $O(D)$ records, then these steps take $O(D^2C)$ time that may be thought of as $O(DN')$, where $DC = O(N')$. We see that step 3 dominates the running time. Overall the running time is $O(b_n N' L \tau)$, where $b_n$ is the average number of records in a block (in step 3), $N'$ is the number of clusters by exact matching, $L$ is the maximum aggregated length of the common attributes of records and $\tau$ is the user-defined threshold value.

### Parallel Algorithm

We observe that the above RLA-CL algorithm has several phases, and almost all of these phases have independent working processes. For example, the distance calculation is done within each block. Therefore processors can perform linkage calculations independent of the others. Some steps are difficult to be parallelized optimally. For them we provide experimentally optimized solutions. Some steps are trivial to parallelize. Here we propose the PRLA-CL (Parallel Record Linkage Algorithm—Complete linkage) algorithm. One processor handles the input, output and collaboration with the other processors and is called the master processor and all the other processors are referred to as slave processors.

**Algorithm 4** PRLA-CL Parallel Record Linkage Algorithm using Complete Linkage Clustering

**Input:** A set of data sets and a configuration file

**Output:** A set of complete linkage clusters

1: procedure PRLA-CL
2: The Master reads data from the input files;
3: The Master broadcasts data;
4: for each processor do
5:   Determine which pairs of data sets should be sorted;
6:   Remove duplicates and merge records;
7: end for
8: The Master collects and merges all exact matched clusters;
9: The Master distributes nearly uniformly representative records to each processor;
10: for each processor do
11:   Create blocks of records sharing the same k-mers;
12: end for
13: The Master collects and merges all blocking information;
14: The Master distributes block lists to all the processors nearly uniformly;
15: for each processor do
16: for each block in block list do
17: Construct a graph where the records are vertices and the connections among them are edges;
18: Connect two vertices if the distance (edit distance is one of the distance calculation methods) between them is at most the user defined threshold value;
19: end for
20: end for
21: The Master accumulates edge lists from each processor;
22: The Master finds connected components using these lists just as we do in Algorithm 3;
23: The Master distributes clusters data uniformly to all the processors;
24: for each processor do
25: Perform complete linkage clustering and post processing (same as in Algorithm 3);
26: end for
27: The Master collects these clusters;
28: The Master merges these clusters with records from initial exact matching clusters;
29: return these complete linkage clusters.

As displayed in Algorithm 4, after receiving data from the master, every processor selects pairs of data sets such that attributes of one data set cover all the attributes of the other data set. Then we accumulate records from each pair. Every processor sorts a specific range of records lexicographically. This range is chosen according to a prefix value of concatenated attributes of each record. If we choose the first 2 characters from each record, there are 676 combinations. If we have \( p \) processors, then every processor can keep track of records starting with \( \frac{676}{p} \) character combinations. The master collects and merges all the exact match records. Then the master chooses a representative from every exact matched group. Then it sends nearly an equal number of records to each of the slave processors. The slave processors generate blocks of records sharing some common \( k \)-mers. The master collects this blocking information. It then sorts blocks according to the number of records they contain. Then the master groups some blocks and aggregates squares of the numbers of all records in that group. The master does this grouping in such a way that all the groups have almost the same aggregate value. Then each processor finds the edge lists. The master collects them and finds connected components. Then the master splits these connected components equally among all the other processors. All of them compute the complete linkage clusters within each component. The master gathers all the clusters and expands every representative record by all of its exact matched records.

Analysis

This parallel algorithm distributes most of the work uniformly across all the processors. Major portions of them have been performed independently. Therefore communication cost is negligible with respect to the computational cost. Some steps have to be explained elaborately.

Step 1 takes \( O(N) \) time to read \( N \) records from \( D \) data sets and broadcast them. We see from the sequential algorithm that some pairs of data sets should be sorted to find duplicate records. In PRLA-CL, every processor determines those pairs of data sets. To compare among
records, we concatenate common attributes of those records. We take the first 3 characters from each concatenated string. There may be $s = 26^3, 10^3$ or $36^3$ divisions of records if the characters are from English alphabet, number or alphanumeric values, respectively. Every processor sorts $s/p$ divisions and removes duplicates by generating exact matching clusters. Although each processor does not get the same amount of records, the overall task is almost the same and the consumed time is really negligible compared to the other computations. Experimental results verify this statement. Therefore, if $s_n$ is the maximum number of records of one division, then step 2 takes $O(s_n/p)$ time which is $O(N/p)$, where $s_n = O(N)$. In step 3 every processor performs blocking on $N/p$ records which uses $O(N'(L - l + 1)/p)$ time. In [20] we see some efficient techniques to distribute blocks among the processors. Step 4 consumes $O(b_pb_p^{-2}L_\tau)$ time, where $b_p$ is the average number of blocks in a processor and $b_{pn}$ is the average number of records in a block. Step 5 is straightforward as the master handles the collected data and finds the connected components in linear time in $N'$ and number of connections, which is $O(N')$. In step 6 the master distributes clusters in the same way it did for blocking in step 4. Every processor gets almost the same amount of workload to find complete linkages among the records. We assume that $C$ is the number of intermediate clusters. Therefore, each processor does work in $O(D'C/p)$ or $O(DN'/p)$ time. We see that the parallel algorithm has been perfectly parallelized. Experimental results show almost linear speed-up.

Results

We have implemented RLA-CL in C++ and PRLA-CL in C++ with MPI library. We deployed them on a HPC cluster having processors of 12 Intel Xeon X5650 Westmere cores and 48 GB RAM.

FEBRL [51, 52] is a popular record linkage system. It generates clusters of very high accuracy. TPA(FCED) [64] achieved a similar accuracy with much less time. From Table 6 of [64] we see that TPA(FCED) took 203 ms in an experiment, whereas FEBRL needed 1284 ms. We outperformed TPA(FCED) by devising a novel RLA algorithm [20]. The implementation attained the same accuracy while being several times faster. The RLA paper integrated and analyzed some experimental results on real and simulated data sets. Those results exposed its efficiency and accuracy in real as well as simulated data sets. Those real data sets contained a very low percentage of errors. RLA algorithm works really fine on real data sets. But yet we see it achieved not more than 98% accuracy for real data sets. Accuracy on simulated data sets varies widely due to a broad ranges of errors. In our experiments we count the possible traps of TPA (FCED) and RLA algorithms and show how RLA-CL finds the expected output. We will also show how blocking information affects its performance. We will evaluate efficiency of record linkage algorithm using complete linkage hierarchical clustering over single linkage clustering. We have employed only simulated data sets, which contain much more errors than normal, to verify our statements of efficiency and accuracy of RLA-CL.

Generation of Simulated Data Sets

We generated three types of synthetic data sets. The first type has a data set of 1 million records. We made 10 copies of this data set. Then we introduced one insertion, deletion, or substitution error in the last name attribute of every record with a 15% probability. This means that around 15% of all the records in a data set have one mismatch from its original record. These data sets have the first name and SSN attributes along with some other attributes. We have taken an equal number of records from each data set in our experiments. If the number of records is 1 million, every data set contributes 100,000 records. SSN is a unique attribute for every record. We compute the accuracy using this attribute. This type is used to compare
performances among TPA(FCED) [64], RLA [20] and RLA-CL implementations. The second type of data sets were generated from the previous 1 million records. We copied this data set two more times. Then we inserted, deleted, or substituted one symbol in the last name of each record. This means that every record has at least one mismatch from its original record. We used four original data sets, and these two data sets three times. The third type is used for analysing different aspects of RLA-CL. The original data set has 1,600,000 records. We generate three copies of this data set. We remove different attributes from each data set. Then we introduce one insertion, deletion, or substitution error in the last name of every record. We analyze how RLA-CL works for different numbers and types of attributes. We have then cloned all of these three data sets.

Sequential Algorithm

We have categorized our experimental results into three sections. The first section shows that RLA-CL outperforms RLA and TPA(FCED) in terms of accuracy and removes the chaining phenomenon. The number of blocks and types of blocking fields affect the running time and accuracy of RLA-CL. We explain them in the second section. In the third section we distribute the running time of RLA-CL and show that it does not take much time than RLA, the best-known algorithm in this category. We have divided the output data into four categories to measure accuracy. Type I includes perfect clusters. Each cluster contains all the records of an individual and does not contain any record from other individuals. Every cluster of Type II has records of only one individual, but does not include all of them. All the records of an individual mixed with some records of the other individuals are included in Type III category. A Type IV cluster has some records from one individual mixed with some records of the other individuals. Here we see that Type I clusters are the most preferred. A Type IV cluster is a truly incorrect cluster. Therefore we prefer more records in Type I category and less records in Type IV category.

Table 1 compares our newly devised RLA-CL algorithm with the previously best-known RLA algorithm as well as TPA(FCED). Number of records ranges from 100 thousands to 1 million across the five data sets. We have used the first name as the blocking field and Social Security Number as the accuracy testing attribute. We have used the edit distance calculation method on the first name and last name attributes. We have set 2 as the threshold value and 3 as the value of \( k \).

From Table 1 we see that RLA-CL takes almost the same time as RLA. RLA and TPA (FCED) produced the same number and types of clusters. But we see TPA(FCED) takes much more time than RLA and RLA-CL. The RLA paper explained the inverse relationship between the multiplicity of exact matched records and the running time of RLA. The RLA-CL algorithm includes RLA as a preprocessing step. After preprocessing is done, the generated clusters are of small size. Therefore complete linkage among the small number of records in every cluster consumes a small amount of time. Even for 1 million records RLA-CL spends only 13 seconds more than RLA. These few seconds do complete linkage clustering and post-processing of all the single linkage clusters. Fig 2 shows this time comparison. These results show that RLA-CL provides almost 100% Type I clusters whereas RLA and TPA(FCED) produce around 96%–98% Type I clusters. If we consider 1,000,000 records of 100,000 individuals, RLA-CL only misses perfect clusters of 241 individuals whereas RLA and TPA(FCED) do not find accurately all the records of 1981 people. This difference occurs because of the chaining problem of single linkage clustering. We have shown this blackType I accuracy comparison in Fig 3.

We have seen four types of accuracy in Table 1. Accuracy can also be calculated in terms of receiver operating characteristics (ROC). For the case of two classes, ROC-based accuracy is defined as (the number of true positives + the number of true negatives)/(the total number of
records). We extend this definition of accuracy to more than two classes as follows. Each cluster is associated with a user who has a majority of records in this cluster. We say that this user owns this cluster. A record in any cluster is labeled as correct if it belongs to the owner of this cluster. Now we compute the accuracy as (the number of records with correct labels)/(the total number of records). Note that this definition of accuracy is a natural extension of ROC-based accuracy to more than two classes.

RLA-CL achieves more than 99.9% accuracy and TPA(FCED) and RLA achieve around 97%—99% accuracy for these data sets (shown in Table 2). Fig 4 also shows these results graphically.

Table 1. A comparison among TPA(FCED), RLA and RLA-CL on simulated data sets (generated with a low error rate).

| No Of Records | Algorithm  | Time  | Type I | Type II | Type III | Type IV |
|---------------|------------|-------|--------|---------|----------|---------|
| 100,000       | TPA(FCED)  | 31.01 | 97.38  | 0.00    | 2.62     | 0.00    |
|               | RLA        | 2.15  | 97.38  | 0.00    | 2.62     | 0.00    |
|               | RLA-CL     | 3.08  | 99.88  | 0.01    | 0.11     | 0.00    |
| 200,000       | TPA(FCED)  | 122.77| 93.76  | 2.86    | 3.26     | 0.12    |
|               | RLA        | 7.77  | 93.76  | 2.86    | 3.26     | 0.12    |
|               | RLA-CL     | 10.4  | 99.05  | 0.73    | 0.20     | 0.02    |
| 400,000       | TPA(FCED)  | 432.5 | 95.88  | 1.84    | 2.19     | 0.09    |
|               | RLA        | 26.56 | 95.88  | 1.84    | 2.19     | 0.09    |
|               | RLA-CL     | 32.15 | 99.45  | 0.44    | 0.10     | 0.01    |
| 600,000       | TPA(FCED)  | 878   | 96.92  | 1.39    | 1.63     | 0.06    |
|               | RLA        | 54.50 | 96.92  | 1.39    | 1.63     | 0.06    |
|               | RLA-CL     | 62.54 | 99.61  | 0.32    | 0.07     | 0.00    |
| 800,000       | TPA(FCED)  | 1503.53| 97.57  | 1.09    | 1.29     | 0.05    |
|               | RLA        | 87.66 | 97.57  | 1.09    | 1.29     | 0.05    |
|               | RLA-CL     | 97.62 | 99.70  | 0.25    | 0.05     | 0.00    |
| 1,000,000     | TPA(FCED)  | 2157.46| 98.02  | 0.89    | 1.05     | 0.04    |
|               | RLA        | 129.54| 98.02  | 0.89    | 1.05     | 0.04    |
|               | RLA-CL     | 141.17| 99.76  | 0.20    | 0.04     | 0.00    |

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Fig 2. A comparison of running times of TPA(FCED), RLA and RLA-CL on simulated data sets (generated with a low error rate).

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We have also included results for the second type of data sets in Table 3. These data sets contain a very high error rate. Even for these data sets, RLA-CL shows almost 100% accuracy in finding perfect clusters.

Sometimes one attribute may be error prone than the others. Blocking on that field produces blocks that may not hold all the records of same individuals. Multiple blocking attributes assure better results. We have explored this issue by employing three different experiments. One uses social security number (SSN) and the last name (LN) as blocking attributes, the second one uses only SSN and the last one uses only LN as the blocking attribute. We have used 5-mer on SSN, a numeric attribute, and 3-mer LN, which contains only English alphabet. Table 4 shows these comparisons in terms of running time and accuracy. We have used 6 data sets where 3 data sets have exact clone so that we could remove half of the records only after the exact matching phase.

Table 2. Computation of accuracy of TPA(FCED), RLA and RLA-CL on simulated data sets (generated with a low error rate).

| No Of Records | Algorithm   | Records With Correct Labels | Accuracy in % |
|---------------|-------------|-----------------------------|---------------|
| 100,000       | TPA(FCED)   | 97880                       | 97.88         |
|               | RLA         | 97880                       | 97.88         |
|               | RLA-CL      | 99949                       | 99.95         |
| 200,000       | TPA(FCED)   | 194910                      | 97.46         |
|               | RLA         | 194910                      | 97.46         |
|               | RLA-CL      | 199836                      | 99.92         |
| 400,000       | TPA(FCED)   | 392930                      | 98.23         |
|               | RLA         | 392930                      | 98.23         |
|               | RLA-CL      | 399835                      | 99.96         |
| 600,000       | TPA(FCED)   | 592100                      | 98.68         |
|               | RLA         | 592100                      | 98.68         |
|               | RLA-CL      | 599835                      | 99.97         |
| 800,000       | TPA(FCED)   | 791680                      | 98.96         |
|               | RLA         | 791680                      | 98.96         |
|               | RLA-CL      | 799836                      | 99.98         |
| 1,000,000     | TPA(FCED)   | 991500                      | 99.15         |
|               | RLA         | 991500                      | 99.15         |
|               | RLA-CL      | 999836                      | 99.98         |
Experiments that do blocking on LN take very little time for a small number of records. But the running time increases rapidly for higher number of records. LN has an average length of 5. Therefore every record on an average goes to 3 blocks. For 100,000 records, 300,000 records are stored in 263 or 17576 blocks. Every block holds around 17 records on an average. But when we have 3,200,000 records, we have 17576 blocks to keep 9,600,000 records. Each block has to store on an average 546 records. We know that the distance calculation occurs among records densely within every block. This is the most time consuming phase of our algorithm. On the other side SSN uses 5-mers for blocking. Therefore every record goes to 5 blocks. For 100,000 records 105 or 100,000 blocks hold 500,000 records, which is 5 per block on an average. For 3,200,000 records this number is 160. But the most compelling reason is that some

| No Of Records | Algorithm | Time  | Type I | Type II | Type III | Type IV |
|---------------|-----------|-------|--------|---------|----------|---------|
| 100,000       | TPA(FCED) | 31.11 | 96.36  | 0.00    | 3.64     | 0.00    |
|               | RLA       | 3.07  | 96.36  | 0.00    | 3.64     | 0.00    |
|               | RLA-CL    | 4.21  | 99.86  | 0.06    | 0.08     | 0.00    |
| 200,000       | TPA(FCED) | 119.62| 92.49  | 2.89    | 4.58     | 0.04    |
|               | RLA       | 11.16 | 92.49  | 2.89    | 4.58     | 0.04    |
|               | RLA-CL    | 14.51 | 99.54  | 0.30    | 0.12     | 0.04    |
| 400,000       | TPA(FCED) | 421.24| 94.83  | 1.91    | 3.23     | 0.03    |
|               | RLA       | 39.15 | 94.83  | 1.91    | 3.23     | 0.03    |
|               | RLA-CL    | 46.67 | 99.69  | 0.23    | 0.06     | 0.02    |
| 600,000       | TPA(FCED) | 898.64| 96.07  | 1.46    | 2.44     | 0.03    |
|               | RLA       | 77.95 | 96.07  | 1.46    | 2.44     | 0.03    |
|               | RLA-CL    | 88.66 | 99.77  | 0.18    | 0.04     | 0.01    |
| 800,000       | TPA(FCED) | 1507.45| 96.91 | 1.16 | 1.91 | 0.02 |
|               | RLA       | 129.79 | 96.91 | 1.16 | 1.91 | 0.02 |
|               | RLA-CL    | 143.19 | 99.82 | 0.15 | 0.03 | 0.00 |
| 1,000,000     | TPA(FCED) | 2171.43| 97.47 | 0.95 | 1.56 | 0.02 |
|               | RLA       | 193.85 | 97.47 | 0.95 | 1.56 | 0.02 |
|               | RLA-CL    | 209.45 | 99.85 | 0.12 | 0.02 | 0.01 |
combinations of letters are more frequent than the others. This makes some blocks much larger than the others. But for numerical values every block is almost equally populated. The time needed for two attributes is the summation of these two attributes. We have depicted this scenario in Fig 5. Blocking attribute has a greater impact on accuracy. Our generated records contain errors either in the SSN or the LN. Therefore many blocks may not be able to hold records of the same individuals. But if we take blocking of two attributes, we get around 100% Type I clusters. Blocking on SSN achieves 90% and blocking on LN gets 81% Type I clusters. SSN has a better performance as each record goes to 6 blocks compared to 3 blocks for the LN attribute. Fig 6 displays the impact of blocking attributes over blackType I accuracy.

Table 5 distributes the running time of RLA-CL when SSN has been used as the blocking field. Exact matching is required to remove exact duplicates. Linear time radix sorting

| No Of Records | Algorithm | Time | Record Category |
|---------------|-----------|------|-----------------|
|               |           | Type I | Type II | Type III | Type IV |
| 100,000       | SSN-LN    | 111.51 | 100,000 | 0        | 0       |
|               | SSN       | 107.71 | 89,866  | 10,134   | 0       |
|               | LN        | 5.56   | 80,476  | 19,524   | 0       |
| 200,000       | SSN-LN    | 252.46 | 199,988 | 4        | 8       |
|               | SSN       | 237.83 | 179,972 | 20,020   | 8       |
|               | LN        | 16.20  | 161,522 | 38,470   | 8       |
| 400,000       | SSN-LN    | 537.83 | 399,952 | 16       | 32      |
|               | SSN       | 480.49 | 359,656 | 40,320   | 24      |
|               | LN        | 48.47  | 322,612 | 77,372   | 16      |
| 800,000       | SSN-LN    | 1064.43| 799,904 | 32       | 64      |
|               | SSN       | 822.92 | 719,474 | 80,472   | 48      |
|               | LN        | 169.08 | 644,204 | 155,758  | 32      |
| 1,600,000     | SSN-LN    | 2657.35| 1,599,832| 56       | 112     |
|               | SSN       | 1912.80| 1,439,608| 160,290  | 96      |
|               | LN        | 622.12 | 1,290,514| 309,424  | 48      |
| 3,200,000     | SSN-LN    | 6422.64| 3,199,676| 96       | 192     |
|               | SSN       | 4261.62| 2,877,986| 321,824  | 152     |
|               | LN        | 2379.88| 2,583,536| 616,340  | 88      |

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Fig 5. A comparison of running time for variations of blocking attributes.

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algorithm does this step efficiently. Approximate clustering is the most time consuming phase that includes linkage calculation steps. We find clusters as connected components with almost no time. These three portions are required for both RLA and RLA-CL. The later one requires some extra time to find complete linkages. We see from the table that this value is negligible compared to approximate cluster time. We have shown these distributions in Fig 7.

Parallel Algorithm

We have run our parallel algorithm on 3.2 million and 6.4 million records blocking on the last name (LN) attribute. Our parallel experiments have been tested on at most 32 cores of 4 nodes, each node having 8 cores.

Fig 8 shows the running time on different number of processors. We get almost linear speedup. These speedups have been drawn in Fig 9. We see that the most time-consuming part is the single linkage calculation among the records within individual blocks. Different blocks have different numbers of records. Even if two blocks have the same number of records, they may need different time as the time needed depends on matching of records as well as record lengths. We have distributed the runtime of PRLA-CL in Table 6. Detailed time distribution of different tasks of parallel RLA such as broadcast time, communication time, time spent by the master, blocking time, merge time, edgelist calculation time, etc. have been described in [20]. In Table 6 we have included communication time, which aggregates broadcast, communication and merge time, exact matching time, approximation clustering time that covers the generation of blocks and calculation of linkage time, finding connected components time and complete linkage time, which includes complete linkage and post-processing time. The first row shows the running time of the same data for the sequential algorithm.

| No of Records | Exact Cl T | Approx Cl T | Conn Comp T | Comp Link T | Total Time |
|---------------|------------|-------------|-------------|-------------|------------|
| 100,000       | 0.77       | 105.34      | 0.01        | 1.59        | 107.71     |
| 200,000       | 1.90       | 232.25      | 0.03        | 3.65        | 237.83     |
| 400,000       | 3.62       | 468.84      | 0.06        | 7.97        | 480.49     |
| 800,000       | 6.36       | 803.65      | 0.08        | 12.83       | 822.92     |
| 1,600,000     | 13.92      | 1870.60     | 0.17        | 28.11       | 1912.80    |
| 3,200,000     | 29.59      | 4172.79     | 0.35        | 58.89       | 4261.62    |

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Fig 7. Running time distribution of RLA-CL.
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Fig 8. Running time of PRLA-CL for 3.2M and 6.4M records.
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Fig 9. Speedup of PRLA-CL for 3.2M and 6.4M records.
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We have achieved 7.5 speedup for 8 cores in a single node, 14.1 for 16 cores across 2 nodes and 26.1 for 32 cores of 4 nodes. Table 6 also shows that communication time is very negligible as most of the steps of the parallel algorithm are easily parallelized. Therefore communication is needed after each phase only. These speedups are great as they are almost linear, but we can improve these speedups if we can ensure a better uniform distribution of blocks among the processors in terms of needed calculation time.

### Discussion

From the results section we see that the single linkage clustering algorithm suffers from the chaining problem. RLA-CL overcomes this problem by employing complete linkage clustering. Accuracy performance of our new algorithm sometimes depends on the number and type of blocking attributes. There is a trade-off between time spent and accuracy. Stable fields should be chosen as blocking attributes. The value of k also affects the running time and accuracy. If we use 4-mers instead of 3-mers, there will be more blocks. Each block will contain less records on an average and therefore it will cost less than before. But a 4-mer creates less substrings of records which will decrease the accuracy. RLA-CL works on different numbers and types of attributes. It does post-processing on complete-linkage clusters based on priority-list attributes. Another major factor that has a great impact on efficiency and accuracy is the threshold value. RLA applies a constant as well as a proportional threshold value and provides those results. Our new algorithm works in the same way. We can apply different threshold values on the training data sets to find out the perfect threshold value for these data sets. There is no universal threshold value for all types of data sets. Errors introduced in the data sets also have an effect on the threshold value and the performance of the algorithms. If the error rate is low, a threshold value of 1 works fine most of the time.

### Conclusions

Our newly developed record linkage algorithms using complete linkage clustering outperform previous best-known algorithms in this category. They produce more accurate results than the others. The exact matching phase sometimes shrinks much-cleaner real data sets a lot by removing duplicate records. Our experiments show that the post-processing phase generates more accurate results. Our parallel algorithm achieves almost linear speedup that can be applied over millions of records on hundreds of processors. Therefore our proposed algorithms, RLA-CL and PRLA-CL provide the best solution for record linkage problems.

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Table 6. Distribution of running time on multiple cores.

| Proc | Comm | Exact | Approx | Conn Comp | Comp Link | Total | Speedup |
|------|------|-------|--------|-----------|-----------|-------|---------|
| Seq  | 0    | 55.31 | 9218.79| 0.68      | 116.19    | 9390.97| -       |
| 1    | 0    | 55.87 | 9219.32| 0.69      | 115.98    | 9391.86| 1.00    |
| 2    | 0.87 | 28.35 | 4655.38| 0.63      | 59.22     | 4744.45| 1.98    |
| 4    | 0.91 | 15.03 | 2403.09| 0.65      | 31.23     | 2450.91| 3.83    |
| 8    | 1.08 | 7.40  | 1226.59| 0.70      | 17.01     | 1252.78| 7.50    |
| 16   | 5.27 | 4.18  | 647.63 | 0.64      | 9.33      | 667.05 | 14.08   |
| 32   | 7.87 | 2.39  | 343.41 | 0.63      | 5.57      | 359.87 | 26.10   |

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Author Contributions
Conceived and designed the experiments: AM SR. Performed the experiments: AM. Analyzed the data: AM SR. Contributed reagents/materials/analysis tools: RA. Wrote the paper: AM SR RA.

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