Temporal group linkage and evolution analysis for census data

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ABSTRACT
The temporal linkage of census data allows the detailed analysis of population-related changes in an area of interest. It should not only link records about the same person but also support the linkage of groups of related persons such as households. In this paper, we thus propose a new approach to both temporal record and group (household) linkage for census data and study its application for change analysis. The approach utilizes the relationships between individuals to determine the similarity of groups and their members within a graph-based method. The approach is also iterative by first identifying high quality matches that are subsequently extended by matches found with less restrictive similarity criteria. A comprehensive evaluation using historical census data from the UK indicates a high effectiveness of the proposed approach. Furthermore, the linkage enables an insightful analysis of household changes determined by so-called evolution patterns.

1. INTRODUCTION
Census data provides valuable information about individuals and households within cities or regions at a specific point in time [18]. Moreover, the temporal linkage of different census datasets allows analyzing the changes that occur in a population which is of increasing importance for social, demographic, economic and health-related studies [8, 13, 18]. In general, the temporal analysis of changing information about individuals and other entities is seen as a major requirement and challenge for future data analysis [6].

There is a large number of available census datasets for different regions of interest. Normally such census datasets are collected on a regular basis, e.g., every ten years, so that multiple successive versions can be utilized to analyze population- and household-related changes. A key prerequisite for such change studies is the temporal linkage of person records as well as of households, representing a group of individuals living together. There has been a modest amount of previous work on such temporal linkage problems, mainly focusing on temporal record linkage taking into account that linkage-relevant attributes such as surname, address or occupation may change over time [2, 5, 15, 17] (see Section 6). These studies mostly ignore the relationships between individuals, e.g., people living together in a household. Moreover, they do not consider the linkage and evolution of groups of related individuals, such as in a household, which is a main focus of this paper.

Fig. 1 illustrates the problem for two successive historical census datasets from 1871 and 1881. In each dataset, individuals are associated to a single household and have a household-specific relationship or role, such as head of household or daughter (of the head of household). These relationships can be represented in household graphs as shown in the lower part of Fig. 1. To understand the changes between the two considered points in time, one has to find matching individuals and their changes which is challenging, in particular due to the occurrence of frequent names (first names like ‘John’ and ‘Elizabeth’ or surnames like ‘Ashworth’ and ‘Smith’ in our dataset) and attribute changes. Of course, we also need to identify people who occur only in one of the datasets because of deaths, emigration, births and immigration. Obviously, a person in one census dataset should match to at most one person in another census dataset so that temporal linkage aims at a 1:1 mapping between person records. Moreover, we want to identify household-related changes, e.g., to what degree the individuals in a household have stayed together or moved to other households. In this case, we have to identify a many-to-many mapping between households.

In our example in Fig. 1, the daughter of the head of household in \( g^{1871}_1 \) (Alice) married Steve from household \( g^{1871}_2 \) and they both moved into the new household \( g^{1881}_1 \) as shown in the 1881 census data (see blue nodes in household graphs). John Riley died within the considered time...
Figure 1: Example census data for two points in time (1871 and 1881). Red / green / blue colored nodes denote individuals who disappear / newly appear / moved to another household.

period (red node for the first census), while the child Mary Smith was born (green node for the second census). Furthermore, a new family (household g_{1881}) moved into the region. Note that the groups g_{1881} and g_{1881} have highly similar attribute values, but only g_{1871} should be linked to g_{1881}. To overcome such ambiguities of person-related attributes, our linkage approach will utilize stable attributes (such as birth year) as well as stable relationships between records, such as family relations or age differences.

In this paper, we propose and evaluate a novel approach for temporal group and record linkage for historical census data that considers the relationships between individuals. Moreover, we use the linked information for an initial change analysis for individuals and households. Specifically, we make the following contributions:

- We propose a new graph-based approach to linking households and person records between successive versions of census data. The approach works in several steps and utilizes an approximate record matching approach to identify pairs of related households. The linkage of households is based on their graph representation, and identifies common subgraphs representing households for the considered census datasets. We then discuss related work and conclude.

2. PROBLEM DEFINITION

Our approaches to temporal linkage and evolution analysis work on a set of census datasets \( \mathcal{D} \) referring to different points in time. Each dataset \( D_i \) of time \( t_i \) consists of a set of person records \( R_i \) and a set of groups \( G_i \) representing households. The records in \( R_i \) are homogeneously structured and have attributes such as first name, surname, age, occupation, and so on. A group \( g_i \in G_i \) consists of associated person records (household members) of \( R_i \) as well as relationships between them. Each record is part of one group (household) only, i.e., groups are not overlapping.

Groups are represented as (household) graphs \( g_i = (V_i, E_i) \) where the vertices of \( V_i \) correspond to the group members and the edges of \( E_i \) represent their relationships. Relationships (edges) have attributes or properties, in particular a relationship type or role, e.g., daughter. Such relationships can be part of the input data (as in Fig. 1) or can be derived later, e.g., the age difference between two persons. For our example, we may record in the graph for group \( g_{1871} \) only the role daughter between Alice and her father John but also the age difference 31 (39-8). Our algorithm not only determines additional properties such as age differences but also additional relationships among group members, e.g., that Alice and William are siblings with an age difference of 6.

In the next section, we formalize our problem of temporal record and group linkage. The linkage approach is described in Section 3, while Section 4 discusses the use of evolution patterns for change analysis. In Section 5, we evaluate our temporal linkage approach and analyze the evolution of households for the considered census datasets. We then discuss related work and conclude.
in \( R_{i+1} \) and vice versa:

\[
M_{R}^{i+1} := \{(r_i, r_{i+1}) | (r_i, r_{i+1}) \in R_i \times R_{i+1} \land \exists (r_i, r'_{i+1}) \in M_R \rightarrow r'_{i+1} = r_{i+1} \land \exists (r'_{i}, r_{i+1}) \in M_R \rightarrow r'_{i} = r_i \}
\]  

(1)

A group mapping \( M_{G}^{i+1} \) consists of group pairs where a group \( g_i \) of \( G_i \) corresponds completely or partially to a group \( g_{i+1} \) of \( G_{i+1} \) according to the common records:

\[
M_{G}^{i+1} := \{(g_i, g_{i+1}) | (g_i, g_{i+1}) \in G_i \times G_{i+1} \}
\]  

(2)

Group mappings can be of cardinality many-to-many (N:M) since persons of a household can match persons of several households in a different census.

For our running example of Fig. 1, the record mapping includes seven person links between the white and blue colored graph vertices, e.g. link \((1871,1,1888,1)\) for John Ashworth and \((1871,3,1888,7)\) for the link between Alice Ashworth and Alice Smith. The two groups in the first census dataset are split among two groups each in the second dataset, so that there are four group links including \((g^r_{1871}, g^r_{1881})\). In our evolution analysis, we will also consider person records and groups that are not reflected in these mappings, e.g. relating to newly occurring or disappeared persons and households.

3. TEMPORAL GROUP LINKAGE

Determining the record and group mappings for the temporal linkage of census datasets is challenging not only due to changing attribute values for the same person (e.g., for surname or occupation) but also due to the high ambiguity and frequent occurrence of certain attribute values, as well as because of data quality issues, e.g., misspelled names, errors for age etc. Group linkage has hardly been studied before \(^1\) and requires a flexible approach to determine many-to-many mappings taking into account that households may split or merge. Similar in spirit to collective entity resolution \(^1\), we determine the similarity between records not only based on attribute values but also considering relationships between records (persons) within a graph-based approach. Furthermore, we not only address record linkage but also solve record and group linkage jointly within a combined approach. To better deal with the partially low similarity of matching person records and the need to determine many-to-many group mappings we propose an iterative approach for temporal linkage. We first identify safe matches with a high similarity and then continuously relax the similarity criterion to find additional records and group links.

Algorithm 1 describes our approach for determining a group mapping \( M_{G}^{i+1} \) and a record mapping \( M_{R}^{i+1} \) between two successive census datasets \( D_i \) and \( D_{i+1} \). The input of the algorithm includes two similarity functions for record matching and parameters for the iterative adjustment of a similarity threshold \( \delta \). We first give a high-level description of the algorithm and its main steps. These steps are then explained in more detail in the four following subsections of this section.

At first, we enrich the graphs for each group (household) in the two input datasets by adding implicit relationships between group members, such as derivable family relations. Moreover, we compute for each relationship between persons the age difference as an additional relationship property for later use in the similarity computations.

The main part of the algorithm is a loop to iteratively identify and extend the group mapping \( M_{G}^{i+1} \) and the record mapping \( M_{R}^{i+1} \). In each iteration, we first apply a similarity function \( Sim_{func} \) to determine an initial linking and clustering of person records based on attribute similarities only (pre-matching step). The similarity function \( Sim_{func} \) specifies the person attributes, a weighting vector \( \omega \), and a similarity threshold \( \delta \) (i.e., two persons are considered to match if the weighted sum of their attribute similarities exceeds \( \delta \)). In the first iteration, we apply a high value \( \delta_{high} \) for \( \delta \) to start with identifying safely matching persons as a basis for also finding safe group matches. Group matches are only determined for pairs of groups connected by at least one (initial) person link. For such group pairs, we apply a subgraph matching to determine shared subgraphs.

**Algorithm 1: Iterative record and group linkage**

**Input:**
- \( D_i \): old census dataset
- \( D_{i+1} \): new census dataset
- \( Sim_{func} \): similarity function for initial record matching
- \( \Delta \): delta for relaxing similarity threshold
- \( \delta_{high} \): upper bound of similarity threshold
- \( \delta_{low} \): lower bound of similarity threshold
- \( Sim_{func,rem} \): similarity function for remaining records

**Output:**
- \( M_{R}^{i+1} \): record mapping
- \( M_{G}^{i+1} \): group mapping

```plaintext
// initialization
1. \( M_{R}^{i+1} \leftarrow \emptyset \) \( M_{G}^{i+1} \leftarrow \emptyset \)
2. \( M_{G}^{12} \leftarrow \emptyset \) \( M_{G}^{12} \leftarrow \emptyset \)
3. \( G_i \leftarrow \text{completeGroups} (G_i) \)
4. \( G_{i+1} \leftarrow \text{completeGroups} (G_{i+1}) \)
5. \( Sim_{func} \leftarrow \delta_{high} \)

// iterative subgraph matching
6. repeat
   // identification of candidates
7. \( C \leftarrow \text{prematching} (R_i, R_{i+1}, Sim_{func}) \)
8. \( \text{subgraph matching and criteria computation} \)
9. \( \text{Sub}_{G_i} \leftarrow \text{subgroups} (C, G_i, G_{i+1}, Sim_{func}) \)
10. \( M_{G}^{i+1} \leftarrow \text{selectGroupMatches} (\text{Sub}_{G_i}) \)
11. \( M_{G}^{i+1} \leftarrow M_{G}^{i+1} \cup M_{G}^{i} \)
12. \( M_{R}^{i+1} \leftarrow \text{recordMapping} (M_{R}^{i+1}, \text{Sub}_{G_i}, R_i, R_{i+1}) \)
13. \( M_{R}^{i+1} \leftarrow M_{R}^{i+1} \cup M_{R}^{i} \)
14. \( M_{R}^{i+1} \leftarrow \text{extract unlinked records and records that are related to unlinked records} \)
15. \( R_i \leftarrow \text{nonMatchedRecords} (R_i, M_{R}^{i+1}) \)
16. \( R_{i+1} \leftarrow \text{nonMatchedRecords} (R_{i+1}, M_{R}^{i+1}) \)
17. \( Sim_{func} \leftarrow Sim_{func} - \Delta \)
18. until \( M_{R}^{i+1} = \emptyset \) or \( Sim_{func} < \delta_{low} \)
19. \( M_{R}^{i+1} \leftarrow \text{match remaining records} \)
20. \( \text{return} < M_{R}^{i+1}, M_{G}^{i+1} > \)
```

---

\(^1\)We are only aware of one approach for group-based linkage of census data \(^8\) that is non-iterative and less sophisticated regarding the use of relationships. In our evaluation in Section 5, we will compare the results for this scheme with our approach.
with both matching persons and matching relationships. In
general, a group of the first census dataset has several can-
didate group matches in the second dataset so that we select
the best group matches considering multiple criteria such as
the degree of record and relationship similarity. The match-
ing subgraphs of linked groups are then used to extract the
matching records for inclusion into the record mapping (line
10 of Algorithm 1).

Further iterations only process records not yet included
in the record mapping determined so far. We continuously
relax the similarity threshold by a decrement $\Delta$ until a
minimal similarity threshold $\delta_{\text{low}}$ is reached (or no fur-
ther group links are identified). Using such relaxed similar-
ity thresholds aims at finding additional matches between
records and groups even in the presence of erroneous or
changed attribute values.

After all iterations are performed we have finished sub-
graph -based group linkage. For the remaining records not
yet associated within matching subgraphs, we apply a sec-
ond attribute-based similarity function $\text{Sim\_func}_{\text{atts}}$ to iden-
tify further person links for inclusion into the record map-
ing (line 17). Moreover, we extend the group mapping by
adding the group pairs that are now linked by the newly
found record links $M_{i,i}^{(r+1)}$ (line 19).

In the following subsections, we describe the discussed
steps in more detail. We start with explaining the pre-
processing step to enrich the existing household graphs by
implicit relationships and additional relationship properties
(Subsection 3.1). In Subsection 3.2, we describe the pre-
matching step of records. In Subsection 3.3, we outline our
subgraph matching approach to identify common subgraphs.
We then introduce the criteria and algorithm used to select
the group matches (Subsection 3.4).

3.1 Group Enrichment

In the initialization phase, we enrich each household group
by adding implicit relationships and stable properties such
as age differences between persons. In our case, each in-
dividual of a household is given a role related to the head
of household (which is a special role). This role may not
be preserved in future census datasets since individuals may
become members of a different household and the head of
household may change as well. Hence, comparing house-
hold graphs by implicit relationships for each record pair
of the original group and replace the head-dependent rela-
tionship types by a unified type. To increase the semantics
of a relationship, we further add the age difference between
two household members as a time-independent relationship
property. Fig. 2 shows an example of the group enrichment
phase for group $g_{1871}$. The relationship between Elizabeth
Smith and Steve Smith is added. Moreover, the age differ-
ences $\text{age\_diff}$ between persons as well as the relationship
types $\text{rel\_type}$ are added to the relationships.

3.2 Pre-Matching

Pre-matching clusters similar records in the census data-
sets based on their attribute similarity and assigns a cluster
label to each record. These labels are utilized to simplify
subgraph matching since the labels identify similar records
without further similarity computation. Pre-matching first applies similarity function $\text{Sim\_func}$ to compare each record of $R_i$ with each record of $R_{i+1}$. The similarity function specifies the attributes to be compared as well as the attribute-specific similarity function, e.g., q-
gram string matching [4]. Furthermore, it uses a weighting vector $\omega$ and a required minimum similarity $\delta$. Applying
the attribute-specific similarity functions to a pair of records $r_i$ and $r_{i+1}$ results is a similarity vector $\text{sim}(r_i, r_{i+1})$. Using $\omega$ we determine an aggregated similarity $\text{agg\_sim}(r_i, r_{i+1})$ by calculating a weighted sum of the attribute similarities:

$$\text{agg\_sim}(r_i, r_{i+1}) = \omega \cdot \text{sim}(r_i, r_{i+1})$$

We then keep only the record pairs whose similarity is above
the specified threshold $\delta$ as potential record matches. Fur-
thermore, we determine the transitive closure or connected
components of these match pairs (record links) to cluster
together all directly and indirectly matching records. We

<table>
<thead>
<tr>
<th>Cluster label</th>
<th>recordID</th>
<th>first name</th>
<th>surname</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>1871_1</td>
<td>john</td>
<td>ashworth</td>
</tr>
<tr>
<td>B</td>
<td>1871_2</td>
<td>elizabeth</td>
<td>ashworth</td>
</tr>
<tr>
<td>C</td>
<td>1871_4</td>
<td>william</td>
<td>ashworth</td>
</tr>
<tr>
<td>D</td>
<td>1871_6</td>
<td>john</td>
<td>smith</td>
</tr>
<tr>
<td>E</td>
<td>1871_7</td>
<td>elizabeth</td>
<td>smith</td>
</tr>
<tr>
<td>F</td>
<td>1871_8</td>
<td>steve</td>
<td>smith</td>
</tr>
<tr>
<td>G</td>
<td>1881_2</td>
<td>mary</td>
<td>smith</td>
</tr>
<tr>
<td>H</td>
<td>1871_5</td>
<td>john</td>
<td>riley</td>
</tr>
<tr>
<td>I</td>
<td>1871_3</td>
<td>alice</td>
<td>ashworth</td>
</tr>
<tr>
<td>K</td>
<td>1881_7</td>
<td>alice</td>
<td>smith</td>
</tr>
</tbody>
</table>

Figure 2: Example of the group enrichment phase
for group $g_{1871}$.  

Figure 3: Pre-matching result for running example. Records
with the same cluster label represent simi-
lar records.
assign to each record of a cluster a unique label, so that records of the same cluster have the same label.

Fig. 3 shows the resulting clusters for the running example by using the attributes first name and surname, \(\omega = (0.5, 0.5)\) and similarity threshold 1. Pre-matching results in the shown ten clusters where all records of a cluster share the same first name and surname. We then assign the cluster labels \(A, B\) etc. to the respective records of the clusters.

### 3.3 Subgraph Matching

Subgraph matching looks for common subgraphs in each pair of groups \(g_i\) and \(g_{i+1}\) of \(G_i \times G_{i+1}\) to determine likely group links. To avoid the computation of the cross product between \(G_i\) and \(G_{i+1}\), subgraph matching is only applied for pairs of groups sharing at least one similar record, i.e., having the same cluster label.

The subgraph \(g_{ub}\) between two groups \(g_i\) and \(g_{i+1}\) (represented by their enriched graphs with \(g_i\) = \((V_i, E_i)\) and \(g_{i+1}\) = \((V_{i+1}, E_{i+1})\)) consists of a set of vertices \(R_{ub}\) and a set of edges \(E_{ub}\). Each vertex in \(R_{ub}\) represents a pair of equally labeled (i.e., similar) records \(v_i\) from \(V_i\) and \(v_{i+1}\) from \(V_{i+1}\). Two vertices \((v_1, v_{1+1})\) and \((v_2, v_{2+1})\) of \(R_{ub}\) are connected by an edge of \(E_{ub}\) if both the old records \(v_1\), \(v_2\) and the new records \(v_{1+1}\), \(v_{2+1}\) of these vertices are connected within their enriched graphs of \(g_i\) and \(g_{i+1}\), respectively.

Furthermore, we require that these edges must have the same relationship type and highly similar relationship properties, in our case regarding the age differences.

Fig. 4 illustrates subgraph matching for group \(g^{1871}_{ub}\) from the first census dataset and the two groups \(g^{1881}_{ub}\) and \(g^{1881}_{ub}\) from the second dataset. For the group pair \((g^{1871}_{ub}, g^{1881}_{ub})\), we have three matching vertices with labels \(A, B\) and \(C\). The three edges have the same relationship types and the same or very similar age differences. The second group pair \((g^{1871}_{ub}, g^{1881}_{ub})\) also shares three vertices with labels \(A, B\) and \(C\) but only one of the edges has the same relationship type and similar age difference. Hence the common subgraph is reduced to the one shown in the bottom right of Fig. 4.

### 3.4 Selection of Group Links

Subgraph matching generates candidates for group linkage based on common subgraphs for different group pairs. There may be several linkage candidates per group in \(G_i\) and in \(G_{i+1}\) so that we have to find the best matching group pairs. The necessary selection should especially guarantee that each record of a group is only linked to one record of another group (This is not the case for the example in Fig. 4 where we have two linkage candidates for members of group \(g^{1871}_{ub}\)). However, a group can link to more than one group if their subgroups are disjoint.

To select for a certain group \(g_i\) the best-matching groups in \(G_{i+1}\) we consider all subgraphs \(g_{ub} = (R_{ub}, E_{ub})\) involving \(g_i\) and apply an aggregated similarity measure. This measure combines three scores capturing the record similarity (Eq. 5), edge similarity (Eq. 6) and the uniqueness (Eq. 7) of a subgroup \(g_{ub}\). The results of the similarity functions are aggregated according to Eq. 4 whereby \(\alpha\) determines the influence of record similarity and \(\beta\) represents the weight of edge similarity.

\[
\text{avg_sim}(g_i, g_{i+1}, g_{ub}) = \frac{1 + 1 + 1}{3} = 1
\]

\[
\text{edge_sim}(g_i^{1871}, g_{ub}^{1881}) = \frac{1 + 1 + 1}{10 + 3} = 0.46
\]

\[
\text{unique}(g_i^{1871}, g_{ub}^{1881}) = \frac{2}{3 + 3 + 3} = 0.66
\]

\[
\text{avg_sim}(g_i^{1871}, g_{ub}^{1881}) = \frac{1 + 1}{2} = 1
\]

\[
\text{edge_sim}(g_i^{1871}, g_{ub}^{1881}) = \frac{1}{10 + 3} = 0.15
\]

\[
\text{unique}(g_i^{1871}, g_{ub}^{1881}) = \frac{2}{3 + 3} = 0.66
\]

For this score we determine the average of the aggregated similarities \(\text{agg_sim}\) for the record pairs of \(R_{ub}\). These aggregated similarities are already determined during pre-matching for each record pair (see section 3.2) and can be obtained from the respective clusters in \(C\).

\[
\text{avg_sim}(g_i, g_{i+1}, g_{ub}) = \frac{\sum_{(r_i, r_{i+1}) \in R_{ub}} \text{agg_sim}(r_i, r_{i+1})}{|R_{ub}|}
\]

- **Average Record Similarity**

- **Edge Similarity**

  The edge similarity \(\text{edge_sim}\) evaluates the similarity of the relationship properties \(\text{rp_sim}\) in the edges in a subgraph, for example the similarity of the age differences between two individuals in the older group \(g_i\) vs. the age difference in the newer group \(g_{i+1}\). Furthermore, we apply an aggregation measure similar to the Dice-Coefficient to relate the edge similarities to the total number of relationships of the considered groups \(g_i\) and \(g_{i+1}\) thereby giving higher weight to those subgraphs covering a large portion of their relationships.

\[
\text{edge_sim}(g_i^{1871}, g_{ub}^{1881}) = \frac{2 \cdot \sum_{e \in E_{ub}} \text{rp_sim}(\text{oldEdge}(e), \text{newEdge}(e))}{|E_i| + |E_{i+1}|}
\]

- **Uniqueness**

  If two group pairs are similar w.r.t both the average record similarity as well as the edge similarity, we like to prefer the group link between the two groups containing records that are less ambiguous than the records of other group pairs. Therefore, we define the uniqueness for a group pair based on the number of vertices of \(R_{ub}\) of \(g_{ub}\) and the aggregated number of records that are assigned to the same label like the records of \(R_{ub}\). The uniqueness is defined as follows:

\[
\text{unique}(g_i, g_{i+1}, g_{ub}) = 2 \cdot \frac{|R_{ub}|}{\sum_{r_i \in R_{ub}} \text{label}(r_i)}
\]
The aggregated similarity of these values reaches a higher value for group pair \((g_{1871}^a,g_{1881}^a)\) than for \((g_{1871}^b,g_{1881}^b)\) due to the higher edge similarity of the former pair. As a result, we would only include group pair \((g_{1871}^a,g_{1881}^a)\) in the group mapping and derive the record mapping only for the common subgraph of this pair.

After the determination of the introduced similarity values per subgroup, we apply Algorithm 2 for the selection of the best-matching group pairs. The algorithm follows a greedy strategy by considering subgraphs in the order of their aggregated similarity score. It also considers the disjointness of subgraphs and can determine group mappings of cardinality N:M.

In each iteration, we select the group pair with the highest group similarity from a priority queue \(pq\). The selected pair \((g_i,g_{i+1})\) is added to the group mapping \(M_G^p\), if the overlap between the already linked records of \(g_i\) as well as \(g_{i+1}\) and the records of the record pairs of \(g_{sub}\) is empty (line 12). Thus, we ensure that a record is linked at most to one record. The linked records are represented by linked\(_R_i\) resp. linked\(_R_{i+1}\). Moreover, the records of \(g_i\) and \(g_{i+1}\) that correspond to a record pair of \(R_{sub}\) of \(g_{sub}\) are represented by the sets \(R^i_{sub}\) and \(R^{i+1}_{sub}\). These sets are returned by getOldRecords and getNewRecords respectively for a certain subgroup \(g_{sub}\). If a group link is added, we update sets of linked records linked\(_R_i\) and linked\(_R_{i+1}\) for \(g_i\) resp. \(g_{i+1}\) (line 14 to 17).

Based on the selected group matches, we are able to identify the record matches contained in the corresponding subgraph \(g_{sub}\). The record links are included in each vertex of \(g_{sub}\) since \(R_{sub}\) is defined as a set of pairs \(r_i\) and \(r_{i+1}\). These pairs are the most appropriate links since the related groups are linked.

### 4. EVOLUTION ANALYSIS

We will now use the results of the temporal record and group linkage to detect changes between different census datasets in order to support the comprehensive evolution analysis of temporal census data. Such a change analysis should not be restricted to a low-level evaluation of individual links but should be realized at a higher, application-specific level to generate relevant and expressive change patterns. We will also include disappearing as well as newly appearing records and groups that are not reflected in the identified mappings but appear only in one of the census datasets. The analysis should further not be limited to two datasets but involve a series of successive census datasets covering longer periods of time.

In this initial study, we use the given census datasets and the determined linkage results to identify a set of basic and more complex changes for records and groups of records that can be identified with the help of so-called evolution patterns (Subsection 4.1). Furthermore, we propose the use of a so-called evolution graph (Subsection 4.2) to provide an aggregated change representation that is extensible to more than two census datasets. Such an evolution graph is a promising basis for advanced graph mining techniques, e.g., to determine frequent or unusual change scenarios.
4.1 Evolution Patterns

We define evolution patterns on individual records and on groups of records. There are three record evolution patterns called preserveR, removeR and addR. We identify these patterns by utilizing the record mapping $M_R^{i+1}$ as well as record sets $R_i$ and $R_{i+1}$ for two successive census datasets $D_i$ and $D_{i+1}$ as follows:

- **preserveR** is a record pair representing one individual linked between $R_i$ and $R_{i+1}$.
  \[ \forall r_i, r_{i+1} \in R_i \times R_{i+1} :\]
  \[ \text{preserve}(r_i, r_{i+1}) \iff \exists (r_i, r_{i+1}) \in M_R^{i+1} \]

- **addR** denotes an individual $r_{i+1}$ in $R_{i+1}$ that is not linked to any record of $R_i$.
  \[ \forall r_{i+1} \in R_{i+1} : \text{add}(r_{i+1}) \iff \exists (r_i, r_{i+1}) \in M_R^{i+1} \]

- **removeR** denotes an individual $r_i$ in $D_i$ that is not linked to any record of $D_{i+1}$.
  \[ \forall r_i \in R_i : \text{remove}(r_i) \iff \exists (r_i, r_{i+1}) \in M_R^{i+1} \]

To analyze the dynamics of groups, we further define group evolution patterns based on changes within groups. These patterns are $\text{add}_G$ and $\text{remove}_G$ as well as the more complex patterns $\text{preserve}_G$, move, split and merge. The patterns $\text{preserve}_G$ and move both relate to pairs of linked groups but differ on whether the linked groups contain at least two preserved members ($\text{preserve}_G$) or only one (move). Each pattern is identified by utilizing the census datasets, the group mapping $M'_G^{i+1}$ and the record mapping $M_R^{i+1}$:

- **addG** denotes a new group $g_{i+1} \in G_{i+1}$ that did not exist in $D_i$. Thus, the group mapping $M'_G^{i+1}$ does not contain any link with $g_{i+1}$.

- **removeG** contains a group of $g_i \in G_i$ that does not exist in $G_{i+1}$ anymore.

- **preserveG** is a group pair connected by a 1:1 link. Moreover, each group of at least 2 individuals satisfying the $\text{preserve}_R$ pattern. This condition allows us to identify preserving households across censuses. The requirement that a ‘preserved’ household should have at least two remaining members is influenced by real-world situations such as households where only the parents remain after their children have moved to another household.

- **move** identifies pairs of linked groups with only one member in common (determined by the $\text{preserve}_R$ pattern) that has moved from the old to the new group (household).

- **split** identifies a change situation between a group $g_i \in D_i$ from the old dataset and a set of groups $g'_1, g'_2, ..., g'_k \in G_{i+1}$ in the new dataset, where at least two individuals of $g_i$ must overlap with each of the groups from $G_{i+1}$. Note, that each individual record can only be contained in one group, i.e., $g'_1, g'_2, ..., g'_k$ are disjoint.

- **merge** covers the opposite situation between a set of groups $g_i, g'_1, ..., g'_k \in G_i$ from the old dataset and one group $g_{i+1} \in G_{i+1}$ from the new dataset, where at least two individuals from groups in $G_i$ must overlap with the merged group $g_{i+1}$. Each individual record can only be contained in one group, i.e., $g_i, g'_1, ..., g'_k$ are disjoint.

Fig. 5(a) shows the corresponding record and group evolution patterns for our running example from Fig. 1. Seven records have been preserved from $D_{1871}$ to $D_{1881}$. Moreover, there are 4 record additions and one removal. According to the defined group evolution patterns, two groups have been preserved (g$^a$ and g$^b$), two groups newly appeared in 1881 ($\text{add}_G$ for g$^a$ and g$^b$) and two persons, Alice (1871,3) and Steve (1871,8), moved from their parents’ households (g$^a_{1871}$ and g$^b_{1871}$) to their own new household g$^a_{1881}$.

4.2 Evolution Graph

Based on the evolution patterns we want to realize further comprehensive evolution analyses for dynamically changing family structures and individual person histories. We propose the use of a so-called evolution graph reflecting the
history of households across two or more successive census datasets. The graph $G_{Evolution}$ captures both the records and groups per census dataset as vertices and interconnects them across successive datasets by edges that are typed according to the identified evolution patterns (change types). Fig. 5(b) shows a sample evolution graph and evolution patterns for two successive versions $D_i$ and $D_{i+1}$. Blue boxes represent group vertices and blue arrows represent group evolution patterns, i.e., the changes between households. Two groups have been preserved and are linked via the group pattern $\text{preserve}_G$ and one household has been split into two households. One individual moved between two households that are thus connected in the evolution graph. The figure also shows the mapping between individual records (gray dotted lines) as well as a new (add$_R$) and a removed (remove$_R$) record without incoming/outgoing edges. The evolution graph enables the application of several graph mining approaches such as cluster analysis, pattern matching or finding frequent subgraphs. One analysis might be to identify households that are preserved across several census periods. A second use case is to identify clusters of related households that can be used for studies of genetic diseases. In Fig. 5(b), a simple computation of connected components on the exemplary evolution graph for two points in time leads to two components consisting of 4 ($CC_1$) and 3 ($CC_2$) households, respectively. Running such a computation for larger households graphs for many successive versions can produce longer chains of connected households, e.g., indicating relationships between many generations of families.

5. EVALUATION

In this section, we evaluate the introduced approaches for temporal record and group linkage for different historical census datasets from the UK that have also been used in a previous study [8]. We first describe these datasets and the evaluation setup in Subsection 5.1. We then evaluate the linkage quality of the new approaches for different configurations (Subsection 5.2). In Subsection 5.3 we compare our approach with the results of the previous study [8] as well as with the collective record linkage approach [14]. Finally, we discuss results of an initial evolution analysis for the considered census datasets.

5.1 Datasets and Setup

In our evaluation, we use six census datasets collected from 1851 to 1901 in ten-year intervals from the district of Rawtenstall in North-East Lancashire in the United Kingdom. Table 1 shows an overview of these datasets according to the number of records and households for the different time periods. The table also shows the number of unique value combinations of the first name and surname attributes to illustrate the degree of ambiguity for these attributes. Furthermore, we report the ratio of missing attribute values. The table shows that the number of households and persons has almost doubled within the 50 years period indicating a substantial population growth. There is a high degree of name ambiguity since each combination of first name and surname is far from unique but has an average frequency of up to 2.23 (for 1851) with a highly skewed frequency distribution due to the presence of frequent surnames such as Ashworth and Smith. Up to 6.5% of the attribute values are missing, which leads to in additional difficulties for finding correct temporal links.

<table>
<thead>
<tr>
<th>Year</th>
<th>1851</th>
<th>1861</th>
<th>1871</th>
<th>1881</th>
<th>1891</th>
<th>1901</th>
</tr>
</thead>
<tbody>
<tr>
<td>$</td>
<td>R_i</td>
<td>$</td>
<td>10334</td>
<td>22429</td>
<td>26229</td>
<td>29051</td>
</tr>
<tr>
<td>$</td>
<td>G_i</td>
<td>$</td>
<td>4070</td>
<td>5576</td>
<td>6025</td>
<td>6374</td>
</tr>
<tr>
<td>$</td>
<td>fn + sn</td>
<td>$</td>
<td>7652</td>
<td>10198</td>
<td>13198</td>
<td>15505</td>
</tr>
<tr>
<td>$\text{ratio}_{mv}$</td>
<td>0.67%</td>
<td>4.19%</td>
<td>3.63%</td>
<td>4.09%</td>
<td>6.33%</td>
<td>6.51%</td>
</tr>
</tbody>
</table>

Table 1: Overview of the census datasets according to the number of records, households, unique combinations of first name and surname $[fn + sn]$ and the ratio of missing values $\text{ratio}_{mv}$.

To evaluate the quality of the group and record mappings in terms of precision, recall and F-measure [4], we use the reference mapping determined in [8]. It covers a subset of 1250 matching households from the 1871 and 1881 datasets that consist of 6864 and 6851 members resp. These households were manually linked by experts by focusing on person records found in both datasets.

In our evaluation, we compare different settings for the similarity function considering the string similarity for five attributes and different weight vectors $\omega$ as shown in Table 2. We also evaluate different similarity thresholds for pre-matching as well as different weights for determining the aggregated group similarity for selecting group links.

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Matching method</th>
<th>$\omega_1$</th>
<th>$\omega_2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>First name</td>
<td>q-gram</td>
<td>0.2</td>
<td>0.4</td>
</tr>
<tr>
<td>Sex</td>
<td>exact</td>
<td>0.2</td>
<td>0.2</td>
</tr>
<tr>
<td>Surname</td>
<td>q-gram</td>
<td>0.2</td>
<td>0.2</td>
</tr>
<tr>
<td>Address</td>
<td>q-gram</td>
<td>0.2</td>
<td>0.1</td>
</tr>
<tr>
<td>Occupation</td>
<td>q-gram</td>
<td>0.2</td>
<td>0.1</td>
</tr>
</tbody>
</table>

Table 2: Compared set of attributes and the corresponding weighting vector $\omega$ to identify the set of blocks $B$ that are used for the subgraph matching.

5.2 Linkage Evaluation

We first analyze the influence of different similarity functions during pre-matching and then discuss the impact of different similarity functions for selecting matching group pairs. Afterwards we study the effectiveness of incremental linkage.

5.2.1 Influence of pre-matching configuration

The proposed linkage approach builds on the initial record matching and clustering performed in the pre-matching step. We thus start our analysis by comparing the results for determining the attribute similarities based on the two weighting schemes $\omega_1$ and $\omega_2$ (Table 2) and different lower similarity threshold bounds $\delta_{low}$. For iterative matching we use a start value $\delta_{high} = 0.7$ for the similarity threshold $\delta$ and $\Delta = 0.05$ for decrementing the threshold until the minimal value $\delta_{low}$ is reached.

Table 3 shows the resulting group and record mapping quality in terms of precision, recall and F-measure for the two weighting schemes and four values of $\delta_{low}$ ranging from 0.4 to 0.55. We observe for all configurations high F-Measure results between 94% and 96% for both the determined record mappings and the group mappings, indicating a very high effectiveness of the proposed approach. The best
<table>
<thead>
<tr>
<th>parameter</th>
<th>(\omega)</th>
<th>(\omega_1)</th>
<th>(\omega_2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\delta_{\text{low}})</td>
<td>0.4</td>
<td>0.45</td>
<td>0.5</td>
</tr>
<tr>
<td>group mapping</td>
<td>Precision (%)</td>
<td>96.1</td>
<td>96.5</td>
</tr>
<tr>
<td></td>
<td>Recall (%)</td>
<td>92.2</td>
<td>92.2</td>
</tr>
<tr>
<td></td>
<td>F-measure (%)</td>
<td>94.1</td>
<td>94.3</td>
</tr>
<tr>
<td>record mapping</td>
<td>Precision (%)</td>
<td>96.6</td>
<td>96.8</td>
</tr>
<tr>
<td></td>
<td>Recall (%)</td>
<td>91.9</td>
<td>91.9</td>
</tr>
<tr>
<td></td>
<td>F-Measure (%)</td>
<td>94.2</td>
<td>94.3</td>
</tr>
</tbody>
</table>

Table 3: Quality of group and record mappings for different weighting vectors \(\omega\) and lower bounds \(\delta_{\text{low}}\).

<table>
<thead>
<tr>
<th>parameter</th>
<th>((\alpha, \beta))</th>
<th>((1.0,0.0))</th>
<th>((0.0,1.0))</th>
<th>((0.5,0.5))</th>
<th>((0.33,0.33))</th>
<th>((0.2,0.7))</th>
</tr>
</thead>
<tbody>
<tr>
<td>group mapping</td>
<td>Precision (%)</td>
<td>92.3</td>
<td>96.7</td>
<td>96.6</td>
<td>96.7</td>
<td>97.3</td>
</tr>
<tr>
<td></td>
<td>Recall (%)</td>
<td>89.1</td>
<td>94.1</td>
<td>94.3</td>
<td>94.4</td>
<td>94.8</td>
</tr>
<tr>
<td></td>
<td>F-Measure (%)</td>
<td>90.7</td>
<td>95.4</td>
<td>95.5</td>
<td>96.0</td>
<td>96.0</td>
</tr>
<tr>
<td>record mapping</td>
<td>Precision (%)</td>
<td>96.2</td>
<td>97.4</td>
<td>97.3</td>
<td>97.3</td>
<td>97.5</td>
</tr>
<tr>
<td></td>
<td>Recall (%)</td>
<td>89.8</td>
<td>93.4</td>
<td>93.4</td>
<td>93.4</td>
<td>93.7</td>
</tr>
<tr>
<td></td>
<td>F-Measure (%)</td>
<td>92.9</td>
<td>95.4</td>
<td>95.3</td>
<td>95.3</td>
<td>95.6</td>
</tr>
</tbody>
</table>

Table 4: Quality of the group and record mappings for different weights \(\alpha\) and \(\beta\) to select matching groups.

F-measure results are generally achieved for \(\delta_{\text{low}} = 0.5\), although the differences are small for the other choices. The simple weighting scheme \(\omega_1\) giving equal weight to each of the five considered attributes is consistently outperformed by the alternate approach giving higher weight to attribute \textit{first name} and only reduced weight for the less stable attributes \textit{address} and \textit{occupation}. Pre-matching with weight vector \(\omega_2\) thus improves F-measure by around 1.7% for the group mapping and up to around 1.3% for the record mapping.

Of course, there are many more possibilities to define the similarity function and we could also apply learning-based methods to find a near-optimal weight vector [4]. Still our results show that using the similarity function with weight vector \(\omega_2\) and \(\delta_{\text{low}} = 0.5\) achieve good and stable results making it an effective default configuration.

### 5.2.2 Similarity weights for selecting matching groups

We now evaluate the influence of the different weights \(\alpha\) and \(\beta\) for determining the aggregated group similarity \(\alpha \cdot \text{sim} + \beta \cdot \text{rel}\), where \(\text{sim}\) driving the selection of matching groups. Table 4 shows the results of the different weights. The quality of the group mapping highly depends on the edge similarity underlining the importance of considering the structural similarity within our household graphs. Without considering the edge similarity \((\beta = 0)\), the F-measure for the group mapping drops to 90.7%, i.e. around 5.3% less than for the best configuration \((\alpha = 0.2, \beta = 0.7)\) and also far less than when ignoring the record similarity \((\alpha = 0)\). The uniqueness score can also improve the overall F-measure. For \((\alpha = 0.2, \beta = 0.7)\) its weight is 0.1 which helped to achieve an improved F-measure compared to the three configurations where it is ignored (when the sum of \(\alpha\) and \(\beta\) equals already 1). The best record mapping is also achieved for \((\alpha = 0.2, \beta = 0.7)\) making it a good default configuration for our datasets.

### 5.2.3 Iterative vs non-iterative linkage

We now want to analyze to what degree the iterative group and record linkage with decreasing similarity thresholds is really helpful compared to a non-iterative, one-shot approach applying only a fixed minimal similarity threshold.

<table>
<thead>
<tr>
<th>method</th>
<th>F-Measure (%)</th>
<th>Recall (%)</th>
<th>Precision (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>group mapping</td>
<td>94.5</td>
<td>94.8</td>
<td></td>
</tr>
<tr>
<td>F-measure (%)</td>
<td>93.8</td>
<td>96.0</td>
<td></td>
</tr>
<tr>
<td>record mapping</td>
<td>91.8</td>
<td>93.7</td>
<td></td>
</tr>
<tr>
<td>F-measure (%)</td>
<td>92.5</td>
<td>95.6</td>
<td></td>
</tr>
</tbody>
</table>

Table 5: Quality of the group mapping and record mapping by using the iterative vs. non-iterative approach.

To evaluate such a non-iterative approach we apply similarity functions with \(\omega_2, \delta_{\text{high}} = 0.5\) and \(\delta_{\text{low}} = 0.5\) resulting in only one iteration. The results are shown in Table 5. We observe that the iterative approach indeed outperforms the non-iterative approach with an F-Measure improvement of \(\approx 2.2\%\) for the group mapping and 3.1% for the record mapping. The improved quality mainly results from a substantially higher precision of more than 97% for both the group and record mapping. This is achieved because the iterative approach finds high-quality matches for the more restrictive thresholds while the more relaxed similarity threshold, with an increased risk of finding wrong matches, is limited to a subset of the records.

### 5.3 Comparison with Existing Approaches

We compare our approach with two previously proposed methods: the collective entity resolution approach of [14] to determine a record mapping as well as the previous group linkage approach [8] for census data. In [14], the authors propose a collective approach that is a specialization of [1]. It initially determines seed record links by applying a high record similarity. The seed links are used to incrementally identify additional links from the neighborhood of the linked records based on their attribute similarity and relational similarity. The overall algorithm follows a greedy strategy that selects in each iteration the record pair with the highest similarity. The related records update their similarities according to the selected record pair. In our implementation, we use the same similarity function as
Table 6: Comparison of our approach with the collective linkage approach of [14] (CL) to determine a record mapping.

<table>
<thead>
<tr>
<th>method</th>
<th>Precision (%)</th>
<th>Recall (%)</th>
<th>F-measure (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CL</td>
<td>93.5</td>
<td>81.2</td>
<td>86.9</td>
</tr>
<tr>
<td>iter-sub</td>
<td>97.5</td>
<td>93.7</td>
<td>95.6</td>
</tr>
</tbody>
</table>

Table 7: Comparison of our approach with the household linkage approach of [8] (GraphSim).

<table>
<thead>
<tr>
<th>method</th>
<th>GraphSim</th>
<th>iter-sub</th>
</tr>
</thead>
<tbody>
<tr>
<td>Precision (%)</td>
<td>97.6</td>
<td>97.3</td>
</tr>
<tr>
<td>Recall (%)</td>
<td>90.1</td>
<td>94.8</td>
</tr>
<tr>
<td>F-measure (%)</td>
<td>93.7</td>
<td>96.0</td>
</tr>
</tbody>
</table>

in our approach (Table 2). Moreover, we filter all record pairs where the normalized age difference is more than 3 years. To generate the seed link, we select the record links with a minimal similarity of 0.9. Table 6 shows the results of the record mapping obtained by collective linking. Our approach outperforms the collective approach w.r.t the record mapping quality by 8.6% for F-measure. The difference between our approach and the collective approach is that we can better link moved records with changed attribute values since we do not only link highly similar records (which is not sufficient for temporal linkage). Furthermore, our subgraph matching utilizes different relationships more comprehensively and benefits from incremental linkage. The previous group linkage approach of [8] initially generates a highly selective record mapping consisting of 1:1 correspondences only. Based on this record mapping, the method calculates an average record similarity and an edge similarity between each group pair. Contrary to our approach, they calculate the similarities based on the initial 1:1 mapping. If correct record pairs are filtered out due to the 1:1 constraint, the approach is not able to identify these links. Hence, this filter step influences the average record similarity as well as the edge similarity, so that correct group links are not identified. Table 7 shows the results of the quality of the group mappings. Our approach achieves a significantly better F-measure for the group mapping compared to [8] (≈3.7%). This improvement is mainly because of a much higher recall that is limited in the previous approach mainly because of the use of the initial 1:1 mapping.

5.4 Analysis of Household Dynamics

Finally, we analyze the evolution of households from 1851 to 1901. For this purpose, we determine the evolution patterns for each successive census dataset pair based on the identified group and record mapping with the best parameter setting. Fig. 6 shows the frequency of each group evolution pattern for each pair of census datasets. In general, we observe an increasing number of households since the number of addG patterns is higher than the number of removeG patterns for each new census. Moreover, we observe an increasing number of preserveG patterns due to the general increase in the number of households over time. From 1891 to 1901, there is also a high number of removeG patterns (up to ≈ 2200) indicating that many households may have moved to a new region. The complex patterns such as split and merge occur only rarely with an average occurrence of ≈ 100 for split and ≈ 70 while the move patterns are more frequent (≈ 1600 on average).

To analyze dependencies between households for the whole time period, we exploit the evolution graph and determine the largest connected component representing all households from 1851 to 1901 that are connected by group patterns. We identified the largest connected component with 17150 households over the complete interval from 1851 to 1901 thereby covering ≈52% of all households. Furthermore, we identify the number of preserved households according to different time intervals for the whole time period from 1851 to 1901. For instance, if we like to identify households that are preserved for 20 years, we define a graph pattern that consists of 2 edges with the pattern type preserveG since the difference between two census datasets is 10 years. Table 8 shows the number of preserved households for the different time intervals. The number of preserving households for all 10 year intervals (1851-61, 1861-71, 1871-81 etc.) represents the overall number of preserveG patterns of the quantitative analysis. Moreover, 260 household are preserved over the whole time period from 1851 to 1901.

6. RELATED WORK

Record linkage or entity resolution has been intensively studied in the past (see [4, 7, 12] for overviews). While the majority of approaches focus on evaluating the similarity of record attributes only, collective or context-based approaches additionally consider the similarity of relationships between entities for improved linkage decisions (e.g. [1, 8, 11, 14, 20, 23]). This idea has also been utilized in our approach but in a tailored way for use within groups such as households. Our approach is especially powerful as it considers different kinds of semantic relationships as well as the
similarity of relationship attributes. Previous collective approaches have also not addressed temporal record linkage in contrast to our scheme.

Relatively few studies have investigated temporal record linkage (e.g., [2, 15, 17]) to link records within dynamically changing data. Existing approaches explicitly consider changing attribute values when matching individual records over time, e.g., by computing value transition probabilities [15]. Temporal clustering approaches as proposed in [3] group temporal records that belong to the same entity to reflect the entity history. Temporal record linkage approaches typically focus on matching individual person records while we also match groups of individuals and identify a record as well as group mapping to interconnect temporal records from census data.

Most closely related to our work is the group-based approach of [8] for matching households in historical census datasets. Our evaluation in Subsection 5.3 has shown that this previous scheme is outperformed by our approach due to its novel features such as an iterative group linkage and subgraph matching based on different semantic relationships. Richards and colleagues investigate in [21] the use of learning-based methods to optimize the use of attribute similarities for temporal record linkage (not group linkage) for census datasets. The observations of this study are complementary to ours and could be used for choosing alternate similarity functions for record matching.

Our work is further related to research on time and evolution-based analysis that is gaining increasing interest. For instance, there are studies analyzing historical web contents to find interesting patterns and trends [25], analyzing person histories on Twitter [16], or collecting and analyzing temporal knowledge from Wikipedia [24]. Our definition of change patterns is further related to previous work in the domain of ontology evolution [10, 22], in particular regarding change detection and diff computation (e.g., [9, 19]). These approaches typically identify basic and complex change operations between different ontology versions. We used this idea to identify time dependent patterns between groups of records to represent the semantics of changes in households over time. Based on the change patterns we are able to realize more comprehensive analysis, e.g., on complex evolution graphs.

7. CONCLUSIONS

We outlined and evaluated a new approach for temporal record and group linkage for the analysis of census data. The approach follows an iterative linkage strategy that first identifies high quality links thereby limiting the more error-prone identification of links between less similar records and groups to subsets of the input data. Group linkage is based on the identification of common subgraphs between groups such as households where we utilize the semantic relationships within groups and relationship properties such as the age differences between individuals. The evaluation showed the high effectiveness of the proposed approach that also outperforms a previous approach for linking census data.

We showed that the linkage results support a detailed evolution analysis of census data at both the level of individuals and groups. We proposed several evolution patterns to identify relevant changes including different kinds of group changes such as splits, merges and the movement of individuals from one group to another. All changes can be maintained within an evolution graph that can be used for a wide spectrum of change analysis, e.g., to identify frequent change patterns or to find connected groups over several census periods.

In future work, we plan to extend the change analysis of census data using the evolution graph and graph mining techniques. We also aim to apply and evaluate the proposed approach on larger census datasets. Furthermore, we want to study additional applications for group linkage, e.g., to analyze the changes in research teams or groups of coauthors over time.

8. ACKNOWLEDGEMENTS

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References


