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INSTITUT NATIONAL DE RECHERCHE EN INFORMATIQUE ET EN AUTOMATIQUE

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## Multi-Dimensional Grid-Based Clustering of Fuzzy Query Results

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**Abstract:** In usual retrieval processes within large databases, the user formulates a first basic (broad) query to target and filter data and next, she starts browsing the answer looking for precise information. We then propose to perform an offline hierarchical grid-based clustering of the data set in order to quickly provide the user with concise, useful and structured answers as a starting point for an online exploration. Every single answer item describes a subset of the queried data in a user-friendly form using linguistic labels, that is to say it represents a concept that exists within the data. Moreover, answers of a given ‘blind’ query are nodes of a classification tree and every subtree rooted by an answer offers a ‘guided tour’ of a data subset to the user. Finally, an experimental study shows that our process is efficient in terms of computational time and achieves high quality clustering schemas of query results

**Key-words:** Database Summary, Flexible Query, Clustering of Query Results

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## Classification Multidimensionnelle des Résultats d'une Requête Floue

**Résumé :** Actuellement, la maturité des SGBDs et leur utilisation intensive dans les systèmes d'information a eu, entre autres, une conséquence majeure dans les entreprises : la production de bases de données volumineuses. La masse d'information ainsi obtenue est difficilement exploitable. En effet, les utilisateurs de ces systèmes doivent faire face au problème de 'surcharge d'information' lors de l'interrogation de leurs données, qui se traduit par un nombre de réponses trop élevé à des requêtes exploratoires. L'organisation et le filtrage de l'information en fonction de l'utilisateur et de la requête est un moyen privilégié pour résoudre ce problème. L'approche proposée dans ces travaux consiste à regrouper les réponses à une requête utilisateur en un ensemble de classes (ou résumés) organisées hiérarchiquement. Chaque classe décrit un ensemble de n-uplets dont les propriétés sont voisines. L'utilisateur pourra ainsi explorer la hiérarchie pour localiser les n-uplets qui l'intéressent et en écarter les autres. Le principe cognitif mis en oeuvre dans cette approche est que les réponses pertinentes sont a priori semblables et donc appartiennent à la même classe d'objets homogènes. Notre algorithme d'interrogation utilise une hiérarchie de résumés (une hiérarchie SaintEtiQ) pré-calculé sur l'ensemble des données pour fournir une classification hiérarchique (collection de résumés) comme réponse à une requête utilisateur. Ainsi, il améliore l'analyse exploratoire par la construction de réponses plus générales, moins nombreuses et mieux ciblées.

**Mots-clés :** Résumé de données, Requetes flexibles, Classification des résultats d'une requête

## 1 Introduction

Because of the ever increasing amount of information stored each day into databases, user can no longer have an exploratory approach for visualizing, querying and analyzing their data without facing the dual issues of Empty and Many answers. In the former case, very few tuples or even no tuples satisfy the user's query, while in the latter, the user is provided with too many tuples forcing her to spend unnecessary time sifting through these items to find relevant results. This is due to the fact that traditional database query processing models have always assumed that the user knows what she wants and consequently tuples that exactly satisfy the selection conditions in the query are returned, no more and no less.

The empty answers problem happens when the user submits a very restrictive query. Many approaches, based for instance on *query relaxation* [1] or *nearest-neighbor* [2] techniques, are proposed to address such problem. These techniques allow the database system to retrieve tuples that closely (though not completely) match the user's query. We do not consider the empty answers problem in this paper.

In contrast, the user faces the so-called many answers (or information overload) problem when she submits a 'broad' query, i.e. , she has a vague and poorly defined information need. Means to circumvent such problem include *query-by-example* [3] [4] and *ranking* [5] [6] based techniques. Such techniques first seek to approximate user's requirement. For instance, systems proposed in [3] [4] use relevance feedback from the user (few records judged to be relevant by the user) while works in [5] [6] use past behavior of the user (derived from available workloads). Then, they compute a score (similarity measures) of every answer that represents the extent to which it is relevant to the estimated user's requirement. Finally, the user is provided with a ranked list, in descending order of relevance, of either all query results or only the top-k [7] subset. The effectiveness of the above approaches depends on their ability to accurately capture the user's requirement, which is, unfortunately, a very difficult and time consuming task.

In this communication, we adopt a completely different strategy to handle many answers problem. It consists in dividing the answer tuples set into different groups, allowing the user to select and browse groups that most closely match what she is looking for, ignoring the irrelevant ones. Indeed, such technique focuses on the user's participation in a cycle of query refinement instead of trying to understand and estimate the user's needs. This idea has traditionally been used for organizing the results of Web search engines [8] [9] [10] [11] but has only recently been used in the context of relational database [12] [13] [14] (see Section 2). However, these approaches are performed on query results and consequently occur at query time. Thus, the overhead time cost is an open critical issue for such *a posteriori* tasks.

To address this issue, we propose to perform an offline hierarchical grid-based clustering of the whole data set in order to quickly provide the user with concise, useful and structured answers as a starting point for an online analysis. Each answer item describes a subset of the queried data in a human-readable form using linguistic labels. Moreover, answers of a given 'blind' query are nodes of the clustering tree and every subtree rooted by an answer offers a 'guided tour' of a data subset to the user. Organizing tuples in nested clusters reduces the

‘information overload’, allowing the user to spend less time sifting and sorting through the results in order to retrieve the required piece of information. The leading idea of this proposal is the *Cluster Hypothesis* [8]: closely associated items tend to be relevant to the same request. However, since such hierarchical grid-based clustering is independent of the query, the set of starting point answers could be large and consequently dissimilarity between items is susceptible to skew. It occurs when the pre-computed clustering schema is not perfectly adapted to the user query. To tackle this problem, we propose an efficient and effective algorithm that rearranges answers based on the hierarchical structure of the pre-computed clustering schema, i.e. , no clustering at all would have to be performed at query time.

The rest of the paper is organized as follows. First, we describe works that relate to our proposal (Section 2). In Section 3 we detail the underlying clustering technique coined SAINTETIQ we consider in this work. Then, in Section 4 we present a querying mechanism to efficiently access SAINTETIQ outputs and we discuss how summary results help facing many answers problem. An algorithm that addresses the problem of cluster dissimilarity by rearranging query results is presented in Section 5. Section 6 discusses an extension of the above process that allows every user to use her own vocabulary when querying the database. An experimental study using real data is presented in Section 7. Finally we give a conclusion and future directions of that work in Section 8.

## 2 Related work

Clustering of query results has been extensively investigated in areas outside database research such as Information Retrieval. The idea was first introduced in the Scatter/Gather system [8], which is based on a variant of the classical K-Means algorithm. It performs clustering of the results after the documents matching the query have been identified by the search engine. Since then, several classes of algorithms have been proposed (SHOC [9], EigenCluster [10], SnakeT [11]). Such post-processing algorithms introduce a noticeable overhead time cost to the query processing, due to the large number of results returned by the search engine. However, they testify that there is a significant potential benefit in providing additional structure in large answer sets.

In contrast to the internet text search, clustering of query results have received much less attention in the database field. An automatic method for categorizing query results (instead of clustering) is proposed in [12]. It performs categorization as post-processing after query results are obtained, with the focus on minimizing the user navigation cost. In [13], authors propose OSQR, an approach for grouping query result based on agglomerative single-link clustering. Given an SQL query as input, OSQR identifies the set of keywords (or values) that occur frequently in the query’s result and then associates each result with a subset of keywords. Next, it clusters keywords based on a keywords distance function. The clustering of keywords, in turn, induces a clustering of the query result rows. In a recent work [14], authors generalize SQL *group-by* operator to enable fuzzy clustering of database query results on a set of attributes specified in the user’s input query. This approach uses a grid-based clustering method based on a variant of the classical K-Means algorithm and generates flat clustering of query results. Note that the number of clusters has to be provided as a

parameter in the query, which assumes that a priori knowledge about the data is available. All the above proposals provide the user with a higher-level view of the data and can be used for an easier navigation in the result. However, categorization and clustering of query results are performed at query time and imply an overhead cost in the query processing.

### 3 Overview of the SAINTETIQ System

In this section, we introduce the main ideas of SAINTETIQ [15], the grid-based clustering model we consider in this work. Then, we give useful definitions and properties regarding our proposal.

#### 3.1 A Two-Step Process

SAINTETIQ takes tabular data as input and produces multi-resolution summaries of records through an online mapping and a summarization processes.

##### 3.1.1 Mapping Service

The SAINTETIQ system relies on Zadeh’s fuzzy set theory [16] and, more specifically on linguistic variables [17] and fuzzy partitions [18] to represent data in a concise form. The fuzzy set theory is used to translate records in accordance with a *Knowledge Base (KB)* provided by a domain expert or even a user. Basically, the operation replaces the original values of every record in the table by a set of linguistic descriptors defined in the *KB*. For instance, with a linguistic variable on attribute *Price* (Figure 1), a value  $t.Price = 95000€$  is mapped to  $\{0.3/cheap, 0.7/reasonable\}$  where 0.3 is a membership grade that tells how well the label *cheap* describes the value 95000. Extending this mapping to all the attributes of a relation could be seen as locating the cells in a grid-based multidimensional space that map records of the original table. The grid is defined by the *KB* and corresponds to the user’s perception of the data space.

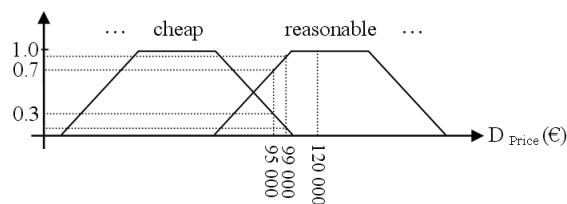


Figure 1: Fuzzy linguistic partition defined on the attribute *Price*

Thus, tuples  $t_1$ ,  $t_2$  and  $t_3$  in Table 1 are mapped into two distinct grid-cells denoted by  $c_1$  and  $c_2$  in Table 2. *old* is a fuzzy label a priori provided by the *KB* on attribute *Age* and it perfectly matches (with degree 1) range  $[19, 24]$  of raw values. Besides, tuple count column gives the proportion of records in  $R$  that belong to the cell and  $0.3/cheap$  says that *cheap* fits the data only with a small degree (0.3). The degree is computed as the maximum of membership grades of tuple values to *cheap* in  $c_1$ .



Table 1: Raw data  $R$ 

Id	Age	Price
$t_1$	22	95000
$t_2$	19	99000
$t_3$	24	120000

Table 2: Grid-cells mapping

Id	Age	Price	tuple count
$c_1$	<i>old</i>	0.3/ <i>cheap</i>	0.4
$c_2$	<i>old</i>	<i>reasonable</i>	2.6

Flexibility in the vocabulary definition of  $KB$  permits to express any single value with more than one fuzzy descriptor and avoid threshold effect due to a smooth transition between two descriptors. Besides,  $KB$  leads to the point where tuples become indistinguishable and then are grouped into grid-cells such that there are finally many more records than cells. Every new (coarser) tuple stores a record count and attribute-dependant measures (min, max, mean, standard deviation, etc.). It is then called a *summary*.

### 3.1.2 Summarization Service

Summarization service is the last and the most sophisticated step of the SAINTETIQ system. It takes *grid-cells* as input and outputs a collection of summaries hierarchically arranged from the most generalized one (the root) to the most specialized ones (the leaves). Summaries are clusters of grid-cells, defining hyperrectangles in the multidimensional space. In the basic process, leaves are grid-cells themselves and the clustering task is performed on  $K$  cells rather than  $N$  tuples ( $K \ll N$ ). From the mapping step, cells are introduced continuously in the hierarchy with a top-down approach inspired of D.H. Fisher's Cobweb, a conceptual clustering algorithm [19]. Then, they are incorporated into best fitting nodes descending the tree. Three more operators could be applied, depending on partition's score  $U$ , that are *create*, *merge* and *split* nodes. They allow developing the tree and updating its current state.  $U$  is a combination of two well-known measures: *typicality* [20] and *contrast* [21]. Those measures maximize within-summary similarity and between-summary dissimilarity. Figure 2 represents the summary hierarchy built from the cells  $c_1$  and  $c_2$ .

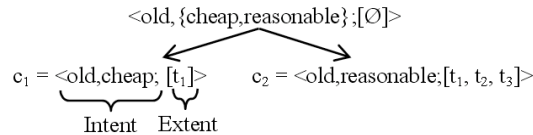


Figure 2: Example of SAINTETIQ hierarchy

### 3.2 Features of Summaries

We introduce in this section basic definitions related to the SAINTETIQ summarization process.

**Definition 1 Summary.** Let  $E = \langle A_1, \dots, A_n \rangle$  be an  $n$ -dimensional space equipped with a fuzzy grid that defines basic  $n$ -dimensional overlapping areas called cells in  $E$ . Let  $R$  be a relation defined on the cartesian product of domains  $D_{A_i}$  of dimensions  $A_i$  in  $E$ . Summary  $z$  of relation  $R$  is the bounding box of the cluster of cells populated by records of  $R$ .

The above definition is constructive since it proposes to build generalized summaries (hyperrectangles) from cells that are specialized ones. In fact, it is equivalent to performing an *addition* on cells:

$$z = c_1 + c_2 + \dots + c_p$$

where  $c_i \in L_z$ , the set of  $p$  cells (summaries) covered by  $z$ .

A summary  $z$  is then an *intensional description* associated with a set of tuples  $R_z$  as its *extent* and a set of cells  $L_z$  that are populated by records of  $R_z$ .

Thus, summaries are areas of  $E$  with hyperrectangle shapes provided by the *KB*. From another point of view, they are nodes of the summary tree built by the SAINTETIQ system.

**Definition 2 Summary Tree** A summary tree over  $R$  is a collection  $\mathcal{Z}$  of summaries verifying:

- $\forall z, z' \in \mathcal{Z}, \quad z \preccurlyeq z' \iff R_z \subseteq R_{z'}$ ;
- $\exists! z \in \mathcal{Z}$  such that  $R_z = \bigcup_{z' \in \mathcal{Z}} R_{z'}$ .

The relation over  $\mathcal{Z}$  (i.e.  $\preccurlyeq$ ) provides a generalization-specialization relationship between summaries. And assuming summaries are hyperrectangles in a multidimensional space, the partial ordering defines *nested summaries* from the larger one to single cells themselves.

Finally, to determine a small set of data representing the whole data set, we need to extract a subset of summaries in the tree. The straightforward way of performing such a task is to define a summary partitioning.

**Definition 3 Summary Partitioning.** The set  $\mathcal{P}$  of leaves of every rooted sub-tree of the summary hierarchy provides a partitioning of relation  $R$ .

We denote by  $P_z$  the top-level partition of  $z$  in the summary tree. It is then the most general partitioning of  $R_z$  we can provide from the tree. Note that the most specialized partitioning is the set of cells covering  $R_z$ , that is  $L_z$ . A partitioning can be obtained *a posteriori* to set the compression rate from the user's needs. For instance, general trends in the data could be identified in the very first levels of the tree whereas precise information has to be looked at near the leaves. Moreover, such a partitioning verifies two basic properties: *disjunction* and *coverage*.

**Property 1 Disjunction**

Summaries  $z$  and  $z'$  are disjoint iff  $\exists k \in [1..n], z.A_k \cap z'.A_k = \emptyset$ .

Regarding such a property, summaries of a partition do not overlap with each other, if we except overlapping of borders of fuzzy cells.

**Property 2** *Coverage*

$$R = \cup_{z \in \mathcal{P}} R_z$$

A partition  $\mathcal{P}$  ensures coverage of the whole data set  $R$  since, by definition, representatives of every branch are included into  $\mathcal{P}$ .

### 3.3 Discussion about SAINTETIQ

The use of SAINTETIQ instead of any other clustering approaches [22] offers a number of advantages. The produced summary hierarchy provides multidimensional views on the data at different levels of granularity through perfectly understandable high-level descriptors. Thus, the user can determine at a glance whether a cluster's content is of interest. Furthermore, SAINTETIQ applies a conceptual clustering algorithm for partitioning the incoming data in an incremental and dynamic way. Thus, changes in the database are reflected through such incremental maintenance of the complete hierarchy (see [15]). Of course, for each application, experts have to be consulted to create linguistic labels as well as the fuzzy-logic membership functions. However, it is worth noticing that, once such knowledge base is defined, the system is fully autonomous with no human intervention required.

As one can observe, the SAINTETIQ hierarchy acts as a multidimensional index. Indexes based on tree structures are widely used for organizing and accelerating access to large multidimensional data sets. The most popular is R-Tree [23] and its variants aR-Tree [24], aP-Tree [25]. R-Tree-based approaches partition the data domain into buckets (i.e., intervals in several dimensions) and build a height-balanced tree that represents the data as nested structured buckets of increasing resolution. Each leaf node is a bucket containing at most  $M$  pointers to the tuples it covers and each inner node contains at most  $M$  pointer to a lower nodes in the tree structure.  $M$  is an input parameter which depends on the hard disk drive's properties (capacity and sector size). These approaches, even though they are very efficient for multidimensional data querying, are not very useful in the context of data discovery, analysis and exploration. Indeed, the grid (buckets) depends on the distribution of the record values rather than the user's perception of the domain. Furthermore, the grouping process depends on the parameter  $M$  and consequently tuples within each group do not meet the cluster hypothesis "closely associated tuples tend to be relevant to the same query".

In the next section, we present a querying mechanism for users to efficiently access the hierarchical summaries produced by SAINTETIQ. Then, we discuss how summary answers reduce information overload that user experiences when querying large databases.

## 4 Querying the SAINTETIQ Summaries

Database systems are being increasingly used for interactive and exploratory data retrieval; examples include users searching their favorite houses, cars, movies, restaurants, and so on over the web. In this context, the user often

has not a specific retrieval goal and is interested in browsing through a set of items (to explore what choices are available) instead of searching among them. For instance, imagine a user who is looking for a house in a database of homes for sale, consisting of a single relation  $R$  with attributes **Price**, **Age**, **Location**. The typical facility available to her is Boolean SQL queries. In terms of query semantics, the user is limited by hard query conditions or constraints (e.g. ,  $\text{Price} \in [150\text{k€}, 250\text{k€}]$ ). However, in many situations, the user actually may not have such hard constraints clearly in her mind. Sometimes it is even impossible to come up with such constraints without knowing what the database contains in the first place. In terms of result organization, what the user gets is a flat table of query results that exactly satisfy all the constraints of the query and thus are equally relevant to the query. While extremely useful for the expert user, the Boolean retrieval model is inadequate for ad-hoc retrieval by exploratory users who cannot articulate the perfect query for their needs - there is usually too many query results returned for a submitted query forcing the user to spend unnecessary time sifting through these items to find relevant results.

To address this problem, we propose a fuzzy retrieval system that (1) uses the SAINTETIQ hierarchy instead of the database to access data; (2) allows users to express vague requirements instead of crisp ones thanks to linguistic labels defined on each attribute domain in the  $KB$ ; (3) provides users with useful and structured answers as a starting point for an exploratory analysis. However, choosing the fuzzy labels from a controlled vocabulary compels the user to adopt a predefined categorization materialized by the fuzzy cells. In Section 6, we deal with user-specific linguistic labels to overcome this pitfall.

To illustrate our proposal, we introduce here a sample data set  $R$  with 30 records ( $t_1 - t_{30}$ ) represented on three attributes: **Price**, **Size** and **Location**. We suppose that  $\{\text{cheap (ch.)}, \text{reasonable (re.)}, \text{expensive (ex.)}, \text{very expensive (vex.)}\}$ ,  $\{\text{small (sm.)}, \text{medium (me.)}, \text{large (la.)}\}$  and  $\{\text{downtown (dw.)}, \text{suburb (su.)}\}$  are sets of linguistic labels defined respectively on attributes **Price**, **Size** and **Location**. Figure 3 shows the summary hierarchy  $H_R$  provided by SAINTETIQ performed on  $R$ .

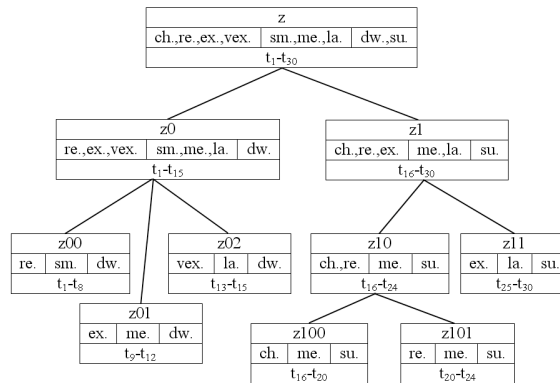


Figure 3: Summary tree  $H_R$  of the data set  $R$

## 4.1 Expression and Evaluation of a Query

The fuzzy querying mechanism proposed here intends to evaluate queries such as Q1:

**Example 1** *Q1: SELECT \* FROM R WHERE Location IN {suburb} AND Size IN {medium OR large}.*

The interpretation we adopted is such that queries are under a conjunctive form: modalities on an attribute are connected with an OR operator and attributes are connected with an AND operator. The AND operator is used because the process should select only data that comply with the characterization on both `Location` and `Size`. On the other hand, the use of the OR operator is due to data constraint: database records are single-valued tuples.

In the following, we denote by  $X$  the set of attributes specified in the user's input query and  $Y$  the set of missing attributes which is the complement of  $X$  relatively to  $R$ :  $X \cup Y = R$  and  $X \cap Y = \emptyset$ . Further, for each attribute  $A \in X$ ,  $Q_A$  denotes the set of fuzzy labels (called *required features*) defined on attribute  $A$ . The set of sets  $Q_A$  is denoted by  $Q^X$  (see example 2).

**Example 2** *Consider query Q1 stated above. We have  $X = \{\text{Location}, \text{Size}\}$ ,  $Y = \{\text{Price}\}$ ,  $Q_{\text{Location}} = \{\text{suburb}\}$ ,  $Q_{\text{Size}} = \{\text{medium}, \text{large}\}$  and  $Q^X = \{Q_{\text{Location}}, Q_{\text{Size}}\}$ .*

Let  $L_A(z)$  be the set of descriptors that appear in a summary  $z$  on attribute  $A$ . When confronting  $z$  with a query  $Q$  to decide whether it corresponds to that query and can then be considered as a result, three cases might occur:

- a:** no correspondence (i.e. ,  $\exists A \in X, L_A(z) \cap Q_A = \emptyset$ ). For one attribute or more,  $z$  has no required feature, i.e. , it shows none of the descriptors mentioned in query  $Q$ .
- b:** exact correspondence (i.e. ,  $\forall A \in X, L_A(z) \subseteq Q_A$ ). The summary  $z$  matches the query  $Q$  semantics. It is considered as a result.
- c:** no decision can be made (i.e. ,  $\exists A \in X, L_A(z) - Q_A \neq \emptyset$ ). There is one attribute  $A$  for which  $z$  exhibits one or many descriptors besides those strictly required (i.e. , those in  $Q_A$ ). Presence of required features in each attribute of  $z$  suggests, but does not guarantee, that results may be found in the subtree rooted by  $z$ . Exploration of the subtree is necessary to retrieve possible results: for each branch, it will end up in situations categorized by case a or case b. Thus, at worst at leaf level, an exploration leads to accepting or rejecting summaries; the problem of indecision is always solved.

The situations stated above reflect a global view of the confrontation of a summary  $z$  with a query  $Q$ .

## 4.2 Search algorithm

The Explore-Select Algorithm (ESA) applies the above matching procedure over the whole set of summaries, organized in a hierarchy, to select relevant summaries. Since the selection should take into account all summaries that

correspond to the query, exploration of the hierarchy is complete. Algorithm 1 describes the ESA algorithm with the following assumptions:

- the function returns a list of summaries;
- function *Corr* symbolizes the matching test reported in Section 4.1;
- operator ‘+’ performs a list concatenation of its arguments;
- function *Add* is the classical constructor for lists, it adds an element to a list of the suitable type;
- $L_{res}$  is a local variable.

---

**Algorithm 1** Function Explore-Select( $z, Q$ )
 

---

```

 $L_{res} \leftarrow \langle \rangle$ 
if Corr( $z, Q$ ) = indecisive then
  for all child node  $z_{child}$  of  $z$  do
     $L_{res} \leftarrow L_{res} + \text{Explore-Select}(z_{child}, Q)$ 
  end for
else
  if Corr( $z, Q$ ) = exact then
    Add( $z, L_{res}$ )
  end if
end if
return  $L_{res}$ 

```

---

The ESA algorithm is based on a depth-first search and relies on a property of the hierarchy: the generalization step in the SAINTETIQ model guarantees that any descriptor that exists in a node of the tree also exists in each parent node. Inversely, a descriptor is absent from a summary’s intent if and only if it is absent from all subnodes of this summary. This property of the hierarchy permits branch cutting as soon as it is known that no result will be found. Depending on the query, only a part of the hierarchy is explored.

Thus, results of Q1 (Example 1), when querying the hierarchy shown in Figure 3, look like:

Table 3: Q1 results

Id_Sum	Price	Size	Location
$z1$	<i>cheap</i> <i>reasonable</i> <i>expensive</i>	<i>medium</i> <i>large</i>	<i>suburb</i>

In this case, the ESA algorithm first confronts  $z$  (root) with Q1. Since no decision can be made, Q1 is respectively confronted to  $z0$  and  $z1$ , the children of  $z$ . The subtree rooted on  $z0$  is then ignored because there is no correspondence between Q1 and  $z0$ . Finally  $z1$  is returned because it exactly matches Q1. Thus, the process tests only 30% of the whole hierarchy.

Note that, the set of records  $R_{z1}$  covered by  $z1$  can be returned (SHOWTUPLES operation) by simply transforming the intent of  $z1$  into an SQL query  $q_{z1}$  and sending it as a usual query to the database system. The WHERE clause of  $q_{z1}$  is generated by transforming the fuzzy labels (vague criteria) contained in the intent of  $z1$  into crisp ones. In other words, each linguistic label  $l$  on an attribute  $A$  is replaced by its support, i.e., the set of all values in  $D_A$  that belong to  $l$  with non-zero membership. Then, the obtained crisp criteria on each summary's attribute are connected with OR operator and summary's attributes are connected with AND operator to generate the WHERE clause. Thus, performing a SHOWTUPLES operation takes advantage of the optimization mechanisms that exist in the database system.

Furthermore, tuples covered by  $z1$  can also be sorted, thanks to their satisfaction degrees to the user's query, using an overall satisfaction degree. We assign to each tuple the degree to which it satisfies the fuzzy criteria of the query. There are many propositions in the literature for defining aggregation connectives [26]. Usually, the minimum and maximum functions stand for the conjunctive and disjunctive connectives.

The ESA algorithm is particularly efficient. In the worst case (exploration of the hierarchy is complete), its cost is given by:

$$C_{ESA}(L) = \varepsilon \cdot \frac{L-1}{w-1} \in O(L)$$

where  $L$  is the number of leaves (cells) of the tree,  $w$  its average width and coefficient  $\varepsilon$  corresponds to the time required for confronting one summary in the hierarchy against the query.

### 4.3 Multi-Scale Summarized Answers

Given a fuzzy query  $Q$ , the ESA algorithm produces a set of clusters (summaries) from a SAINETIQ hierarchy instead of a list of tuples  $tset(Q)$ . Each answer item  $z$  describes a subset of the query result set  $tset(Q)$ . Hence, the user can unambiguously determine whether  $R_z$  contains relevant tuples only by looking at the intensional description of  $z$  and particularly on unspecified attributes ( $Y$ ) because all answer tuples satisfy the specified conditions related to attributes in  $X$ . Three cases might occur:

- 1:** summary  $z$  doesn't fit user's need. It means that for at least one unspecified attribute  $A \in Y$ , all linguistic labels (the set  $L_A(z)$ ) are irrelevant to the user. For instance, consider the Q1 results (Table 3).  $z1$  doesn't contain any relevant tuple if the user is actually looking for *very cheap* homes because *very cheap*  $\notin L_{Price}(z1)$ . In other words, none of the records in  $R_{z1}$  is mapped to *very cheap* on attribute **Price** and consequently a new broad query with less selective conditions may be submitted or the task may be abandoned (we denote this with the IGNORE option).
- 2:** summary  $z$  exactly fits user's need. It means that for each  $A \in Y$ , all linguistic labels in  $L_A(z)$  are relevant to the user. Assume that the user is interested in *cheap*, *reasonable* as well as *expensive* homes. Thus, all tuples contained in  $R_{z1}$  are relevant to her. In such cases, she uses SHOWTUPLES option to access tuples stored in  $R_z$ . The SHOWTUPLES option is similar to the one defined by Chakrabarti et al. in [12].

- 3:** summary  $z$  partially fit user's need. In this case, there is at least one attribute  $A \in Y$  for which  $L_A(z)$  exhibits too many linguistic labels regarding the user's requirement. For instance, the set  $R_{z1}$  partially matches the needs of a user who is looking for *cheap* as well as *reasonable* homes because  $R_{z1}$  contains also tuples that are mapped to *expensive* on attribute *Price*. In this case, a new query with more selective conditions (e.g. , *Price IN{cheap OR reasonable}*) may be submitted or a new clustering schema of the set  $R_z$ , i.e. , which allows to examine more precisely the dataset, is required. Since  $z$  is a hyperrectangle, we present to the user the children of  $z$  (SHOWCAT option). Each child of  $z$  represents only a portion of tuples in  $R_z$  and gives a more precise representation of tuples it contains. For example,  $\{z_{10}, z_{11}\}$  partitions  $R_{z1}$  into two subsets  $R_{z_{10}}$  and  $R_{z_{11}}$ ;  $z_{10}$  exactly fit user needs. Since the entire tree is pre-computed, no clustering at all would have to be performed at feedback time. The SHOWCAT option is similar to the one defined in [12].

More generally, a set of summaries  $S = \{z_1 \dots z_m\}$  (clusters) is presented to the user as a clustering schema of the query result  $tset(Q)$ . The three options IGNORE (case 1), SHOWTUPLES (case 2) and SHOWCAT (case 3) give the user the ability to browse through the  $S$  structure (generally a set of rooted subtrees), exploring different datasets in the query results and looking for relevant information. Indeed, the user navigates through the set  $S = \{z_1 \dots z_m\}$  using the following exploration model:

- i) start the exploration by examining the intensional description of  $z_i \in S$  (initially  $i = 0$ );
- ii) if case 1, ignore  $z_i$  and examine the next cluster in  $S$ , i.e. ,  $z_{i+1}$ ;
- iii) if case 2, navigate through tuples of  $R_{z_i}$  to extract every relevant tuple and thereafter, go ahead and examine  $z_{i+1}$ ;
- iiii) if case 3, navigate through children of  $z_i$ , i.e. , repeat from step (i) with  $S$ , the set of children of  $z_i$ . More precisely, examine the intensional description of each child of  $z_i$  starting from the first one and recursively decide to ignore it or examine it (SHOWTUPLES to extract relevant tuples or SHOWCAT option for further expansion). At the end of the exploration of the children of  $z_i$ , go ahead and examine  $z_{i+1}$ .

This process stops when the user reaches the relevant cluster, i.e. , the one that meets her requirement. Indeed, our general assumption is that relevant tuples for the same user's requirement are more similar (close) to each other and consequently they belong to the same cluster at a given granularity level.

For instance, suppose a user is looking for *medium*, *large* as well as *cheap* homes but issues the query Q1 (Example in Section 1). The set of summaries  $S$  presented to such user is  $\{z1\}$  (Table 3), where  $z1$  is a subtree in the pre-computed hierarchy shown in Figure 3. Thus, the user can explore the subtree rooted by  $z1$  as follows to reach relevant tuples: analyze  $z1$  intent and explore it using SHOWCAT option, analyze  $z_{10}$  intent and explore it using SHOWCAT option, analyze  $z_{110}$  intent and use SHOWTUPLES option to navigate through the tuples in  $R_{z_{110}} = t_{16} - t_{20}$  and identify every relevant tuple, analyze  $z_{101}$  intent and ignore it, analyze  $z_{11}$  intent and ignore it.



Note that when the set  $S = \{z\}$  is a singleton, i.e.,  $z$  is a node of the pre-computed clustering tree, its exploration is straightforward. Indeed, given a summary of the tree rooted by  $z$  the user wishes to examine more closely (SHOWCAT option), its children are well separated since SAINTETIQ is designed for converging to the locally optimized hierarchy. Furthermore, the number of clusters presented to the user, at each time, is small; the highest value is equal to the maximum width of the pre-computed tree. However,  $S$  is often a set of summaries of the pre-computed tree. Since the hierarchical grid-based clustering schema is independent of the query, the size of  $S$  could be large and consequently, dissimilarity between its items is susceptible to skew. It occurs when the pre-computed clustering schema is not adapted to the user query. In such cases, choosing the relevant summaries from the large starting point results  $S$  is a difficult endeavor. In the following section, we describe an algorithm for rearranging query results.

## 5 Rearranging query results

Consider the pre-computed hierarchy  $H$  shown in Figure 4 (Left) and a user's query  $Q$ . Furthermore, suppose that the ESA algorithm returns the set of summaries  $S = \{z_{00}, z_{01}, z_{1000}, z_{101}, z_{11}\}$  when querying  $H$  using  $Q$ . Results for  $Q$  are scattered over the tree. Users often face the above problem since the hierarchical grid-based clustering is independent of the query.

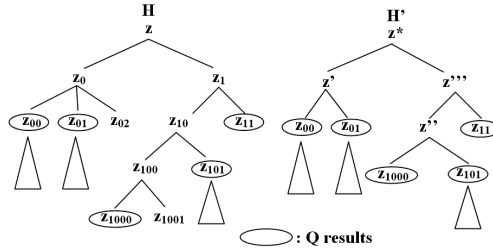


Figure 4: (Left) A pre-computed hierarchy  $H$  (Right) Rearranging  $Q$  summary results

A straightforward way to address this problem would be to, first, execute SAINTETIQ summarization service (SEQ) on the cells populated by records of  $tset(Q)$ . Then, we present to the user the top-level partition of the result tree. Thus, size reduction and discrimination between items in  $S$  are clearly achieved at the expense of an overhead computational cost. Indeed, the SEQ process time complexity is in  $O(L' \cdot \log(L'))$  with  $L'$  the number of cells populated by answer records:

$$C_{SEQ} = k \cdot \log_{w'}(L') \cdot L'$$

where  $w'$  is the average width of the result tree,  $\log_{w'}(L')$  an estimation of its average depth and coefficient  $k$  corresponds to the time required for evaluating the quality of a partition  $P_z$  resulting from the incorporation of a new cell into each of the children nodes of  $z$  as well as additional operations (splitting, merging and creating a new node: see Section 3.1.2).

Thus, the global cost  $C_{ESA+SEQ}(L)$  of such a post-processing algorithm (including search-ESA and summarization-SEQ steps) is in  $O(L \cdot \log(L))$  where  $L$  is the number of leaves of the pre-computed tree ( $L \simeq L'$  since the query is expected to be broad). This solution doesn't fit querying process requirements (see Section 7), that is to quickly provide to the user concise and structured answers.

To tackle this problem, we propose an algorithm, called Explore-Select-Rearrange Algorithm (ESRA), that rearranges answers based on the hierarchical structure of the pre-computed clustering schema. The main idea of this approach is rather simple. It starts from the summary partition  $S$  (a clustering schema of the query results) and produces a sequence of clustering schemas with a decreasing number of clusters at each step. Each clustering schema produced at each step results from the previous one by merging the 'closest' clusters into a single one. Similar clusters are identified thanks to the hierarchical structure of the pre-computed clustering schema. Intuitively, summaries which are closely related have a common ancestor lower in the hierarchy, whereas the common ancestor of unrelated summaries is near the root. This process stops when it reaches a single hyperrectangle (the root  $z^*$ ). Then, we present to the user the top-level partition (i.e., children of  $z^*$ ) in the obtained tree instead of  $S$ .

For instance, when the ESRA process is performed on the set of summaries  $S = \{z_{00}, z_{01}, z_{1000}, z_{101}, z_{11}\}$  stated above, the following sequence of clustering schemas is produced:

$z_{00}$	$z_{01}$	$z_{1000}$	$z_{101}$	$z_{11}$
$z_{00} + z_{01}$		$z_{1000} + z_{101}$		$z_{11}$
$z_{00} + z_{01}$		$z_{1000} + z_{101} + z_{11}$		
$z_{00} + z_{01} + z_{1000} + z_{101} + z_{11}$				

The hierarchy  $H'$  obtained from the set of query results  $S$  is shown in Figure 4 (Right) where  $z' = z_0 - z_{02}$ ,  $z'' = z_{10} - z_{1001}$  and  $z''' = z_1 - z_{1001}$ . Thus, the partition  $\{z', z'''\}$  is presented to the user instead of  $S$ . This partition has a small size and defines well separated clusters. Indeed, all agglomerative methods, among them the ESRA algorithm, have a monotonicity property: the dissimilarity between the merged clusters is monotonically increasing with the level. It means, the dissimilarity value of the partition  $\{z', z'''\}$  is greater than the dissimilarity value of the partition  $\{z_{00}, z_{01}, z_{1000}, z_{101}, z_{11}\}$ .

Algorithm 2 describes the ESRA process. It is a modified version of algorithm 1 with the following new assumptions:

- it returns a summary ( $z^*$ ) rather than a list of summaries;
- function `AddChild` appends a node to caller's children;
- function `NumberOfChildren` returns the number of caller's children;
- function `uniqueChild` returns the unique child of the caller;
- function `BuildIntent` builds caller's intent (hyperrectangles) from intents of its children;
- $Zres$  and  $Z'$  are local variables of type summary.

**Algorithm 2** Function Explore-Select-Rearrange( $z, Q$ )

---

```

Zres ← Null
Z' ← Null
if Corr( $z, Q$ ) = indecisive then
  for all child node  $z_{child}$  of  $z$  do
     $Z' \leftarrow \text{Explore} - \text{Select} - \text{Rearrange}(z_{child}, Q)$ 
    if  $Z' \neq \text{Null}$  then
      Zres.addChild (Z')
    end if
  end for
if Zres.NumberofChildren() > 1 then
  Zres.BuildIntent()
else
  Zres ← Zres.uniqueChild()
end if
else
  if Corr( $z, Q$ ) = exact then
    Zres ←  $z$ 
  end if
end if
return Zres

```

---

This ESRA process (Algorithm 2) returns a summary tree instead of a summary list (Algorithm 1). Such summary tree is built at the same time the pre-computed clustering schema is being scanned. It means that no clustering at all would have to be performed at query time. Hence, the computational complexity of the ESRA algorithm is in the same order of magnitude (i.e.,  $O(L)$ ) than the ESA Algorithm (see Section 4.2):

$$C_{ESRA}(L) = (\gamma + \varepsilon) \cdot \frac{L - 1}{w - 1}$$

where coefficient  $\gamma$  is the time cost for the additional operations (i.e., addChild, uniqueChild and BuildIntent).

The ESRA cost is only a small constant factor  $\gamma$  larger than that of the ESA algorithm since ESRA algorithm takes advantage of the hierarchical structure of the SAINTETIQ summaries to merge similar clusters. However, a limitation relies in the fact that ESRA algorithm restrains the user to queries using a controlled vocabulary (i.e., a summary vocabulary). Indeed, consider a user looking for *cheap* homes. What is meant by *cheap* can vary from one user to another one, or from one kind of users to another one (e.g., *cheap* does not have the same meaning for home buyers and for apartment tenants). In the following, we deal with user-specific linguistic labels to overcome this problem.

## 6 Discussion about Extension to any Fuzzy Predicate

We aim at allowing the user to formulate conditions in selection queries in her own language. As mentioned above, our current querying process cannot handle

free vocabulary. Regarding the query cost and accuracy, the ideal solution to address such problem consists in building a summary tree for every user such that queries and summaries share the same vocabulary (the grid). Thus, the ESRA algorithm, described in Section 5, can be used in its current state. But maintaining *user-specific summaries* is hardly conceivable in a system with multiple users. Two alternatives are then envisaged.

One could consider *group profiles* in order to reduce the number of managed summaries (e.g. , one for buyers and the other for tenants). In this case, the ESRA Algorithm can also be used directly since users formulate queries within the vocabulary of their groups. In this approach, users would subscribe to groups that share the same (or similar) vocabulary as theirs. In addition, they have to be familiar with their group's terms before using them accurately. But this option is not much more convenient than using ad-hoc fuzzy labels, predetermined by a domain expert or even the database administrator. Moreover, it only transposes the problem of user-specific summaries maintenance to group-specific ones.

Probably the most interesting solution consists in building only one SALETTE-TIQ summary tree using an ad-hoc vocabulary, and querying these summaries with user-specific fuzzy labels. In the following, we assume that one summary tree of the database is considered. It is created using a priori defined linguistic variables that constitute the summary vocabulary. Nevertheless, the user defines her own fuzzy labels to formulate free fuzzy predicates within selection queries and then search the summary tree thanks to a vocabulary mapping step.

## 6.1 Query rewriting

Since the vocabulary of the query predicates is different from the one in the summaries, the query must be rewritten. Rewriting query is a fuzzy set-based mapping operation to translate query predicates from the user-specific vocabulary to the summary language (the grid). Consider a fuzzy query expressed using user-specific linguistic labels. Each feature  $F$  (elementary predicate) on an attribute domain  $D_A$  of the query is translated into the smallest superset of linguistic labels  $G$  taken from the summary vocabulary, using the usual fuzzy intersection:

$$F \cap G \neq \emptyset \quad \text{iff} \quad \exists x \in D_A, \min(\mu_F(x), \mu_G(x)) > 0 .$$

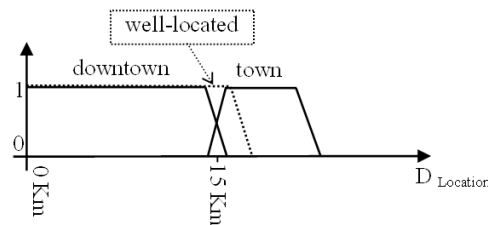


Figure 5: The user's term *well-located* (dashed line) is rewritten with the terms *downtown* and *town*

For instance, in the query “find *well-located* homes”, the user specific term *well-located* is rewritten with the two fuzzy labels of the summaries *downtown* and *town* (Figure 5).

In this scenario, the initial  $Q$  query has been rewritten in a *conservative* manner so that ‘nothing is lost’. Thus, an empty set of summaries  $S = \emptyset$  as an answer guarantees that there is really no result tuples to the query (i.e. ,  $tset(Q) = \emptyset$ ). A non-empty set  $S = \{z_1 \dots z_m\}$  denotes the possibility that answer tuples exist, but nothing can be guaranteed (i.e. ,  $tset(Q) \subseteq \bigcup_{1 \leq i \leq m} R_{z_i}$ ). Of course tuples of  $\bigcup_{1 \leq i \leq m} R_{z_i} \setminus tset(Q)$  do not satisfy the initial query  $Q$ , however, they are close to the initial user’s query and consequently interesting since the user is in an exploratory analysis and tries to discover the content of the database.

In some cases, it could be more suitable to guarantee that answers exist instead of guaranteeing empty answers only. To achieve this, the initial query has to be rewritten using subsets of linguistic labels, using the usual fuzzy inclusion:

$$F \subseteq G \quad \text{iff} \quad \forall x \in D_A, \mu_F(x) \leq \mu_G(x) .$$

For instance, the term *well-located* is only mapped to *downtown* (Figure 5). In this second rewriting scenario, a non-empty set  $S = \{z_1 \dots z_m\}$  as an answer guarantees that all tuples covered by its summaries are really in the result of the query (i.e. ,  $\bigcup_{1 \leq i \leq m} R_{z_i} \subseteq tset(Q)$ ) but there could be others (i.e. ,  $tset(Q) \setminus \bigcup_{1 \leq i \leq m} R_{z_i}$ ) that satisfy predicates and do not belong to  $S$ .

Note that both scenarios can be used together for the same query, as one guarantees null answers and the other guarantees non-null answers. This dual approach can be viewed as reasoning with both Possibility and Necessity (see [27]) of the query result. Moreover, these two scenarios are the boundaries of a spectrum of possible strategies: the generalized expression of the rewriting process consists in defining a threshold  $\tau$  to decide whether the mapping of two fuzzy labels is correct. The user’s label  $d$  is mapped to a summary label  $l$  if and only if the similarity of  $d$  to  $l$ , denoted by  $\sigma(d, l)$ , is greater than or equal to  $\tau$ . The measure  $\sigma(d, l)$  evaluates the satisfiability [28] of a summary fuzzy set  $l$  to a user’s fuzzy set  $d$ . In this work, we use the following definition:

$$\sigma(d, l) = M(d \cap l) / M(l)$$

Assuming the usual definition of the fuzzy intersection  $\mu_{l \cap d} = \min(\mu_l, \mu_d)$  and the following fuzzy measure  $M$ :

$$M(l) = \begin{cases} \sum_{x \in D_A} \mu_l(x) & \text{if } D_A \text{ is finite} \\ \int_{D_A} \mu_l(x).dx & \text{otherwise.} \end{cases}$$

Note that when  $\tau = 0^+$  (respectively  $\tau = 1$ ), we fall back into the Possibility (respectively Necessity) scenario.

Once the query is rewritten, the ESRA algorithm, could be performed in order to provide the summarized answer. Relevance of summaries in the result depends on the query mapping and especially the satisfiability degree of the rewritten predicates. Thus, we propose in the following to sort the query results using their satisfaction to the user’s query.

## 6.2 Sorting the results

Consider an initial user's query  $Q$ , expressed using her own vocabulary: “find all results which are  $d$  or  $d'$  on  $A_i$  and  $d''$  on  $A_j$ ”. The set of attributes for which required features exist is  $X = \{A_i, A_j\}$  and the representation  $Q^X$  of  $Q$  is:

$$Q^X = \{Q_{A_i}, Q_{A_j}\} \text{ where } Q_{A_i} = \{d, d'\} \text{ and } Q_{A_j} = \{d''\}$$

The querying process is performed using  $Q'$ , the rewritten version of  $Q$ . Each summary  $z$  from  $Q'$  results is represented by a set of descriptors in  $L_{A_i}(z)$  on each attribute  $A_i$ . For instance,  $L_{A_i}(z) = \{l, l'\}$ . Once we have the query and the answers represented by linguistic descriptors, we can define a scoring function based on *degree of matching* between the summary and query predicates, using fuzzy set operations. This choice is justified since fuzzy sets can be given a semantic in terms of degree of satisfiability. Indeed, we use  $\sigma(d, l)$  to compute the degree of matching of each query required feature ( $d$ ) from  $Q_{A_i}$  and each result description ( $l$ ) from  $L_{A_i}(z)$ . Thus, the global degree of matching  $\sigma_G(Q_{A_i}, L_{A_i}(z))$  between  $L_{A_i}(z)$  and  $Q_{A_i}$  is a combination of all the pairwise degrees. The simplest way to combine these degrees is by using a disjunctive aggregation that aggregates values as an OR operator, so that the result of the combination is high if some (at least one) values are high. Disjunctive combination is typically performed in fuzzy set theory by T-conorm operators, whose most used instance is 'max'. Thus, we obtain:

$$\sigma_G(Q_{A_i}, L_{A_i}(z)) = \max_{l \in L_{A_i}(z), d \in Q_{A_i}} \sigma(d, l)$$

The function  $\sigma_G$  establishes a comparison, attribute by attribute. Thus, we obtain as many values of comparison as there exist attributes with required features. A last aggregation step of those values is then necessary. The simplest way to do this is by using a conjunctive operator, where we require that each of the result descriptions matches the corresponding part in the query. For instance, we use the product operator to evaluate the degree of satisfiability between a given query  $Q$  and each answer  $z$ . We compute a  $score(Q, z)$  of each answer  $z$  which represents the degree to which  $z$  is relevant for the user query  $Q$ . Such a score is defined as follows:

$$score(Q, z) = \prod_{A_i \in X} \sigma_G(Q_{A_i}, L_{A_i}(z))$$

Thus, for each answer  $z$  from  $S$  (the set of  $Q'$  results returned by ESRA), we compute its score  $score(Q, z)$ ; then we sort these results by their score in decreasing order; and finally return the fully ordered list  $S_{sorted}$  to the user. Furthermore, when a user decides to explore a summary  $z'$  of  $S_{sorted}$  using the SHOWCAT option, we sort the set of children of  $z'$  according to their scores in decreasing order. Note that all basic needed values  $\sigma(d, l)$  are available because  $z'$  intent includes the intent of its children. Likewise, when a user decides to explore a summary  $z'$  of  $S_{sorted}$  using SHOWTUPLES option, each tuple  $t$  can be sorted by  $score(Q, t)$ . Indeed, if  $t$  takes value  $x$  on attribute  $A_i$ ,  $L_{A_i}(t) = \{x\}$  is the fuzzy set whose membership function is defined as follows:

$$\forall y \in D_{A_i}, \mu_x(y) = \begin{cases} 1 & \text{if } y = x; \\ 0 & \text{otherwise.} \end{cases}$$

## 7 Experimental Results

Evaluating and comparing the effectiveness of different approaches that address the Many answers problem in databases is challenging. Unlike Information Retrieval where there exist extensive user studies and available benchmarks (e.g. , the TREC<sup>1</sup> collection), such infrastructures are not available today in the context of Relational Databases. Nonetheless, in this section, we discuss the efficiency and the effectiveness of ESA, ESA+SEQ and ESRA processes based on a real database.

### 7.1 Data Set

Through an agreement, the CIC Banking Group provided us with an extract of statistical data used for behavioral studies of customers. The dataset is a collection of 33735 customers defined on 70 attributes (e.g. , age, income, occupation, etc.). For the results presented in this paper, we used a subset of 10 of the most relevant attributes. On each attribute, marketing experts defined between 3 and 8 linguistic labels leading to a total of 1036800 possible descriptor combinations (i.e. , 1036800 possible cells). Note that all the experiments were done on a 1.7GHz P4-based computer with 768MB memory.

### 7.2 Results

In this section, we validate our process (ESRA) for clustering query results with regard to performance (computational time) and usability. All experiments reported in this section were conducted on a workload composed of 150 queries with a random number of selection predicates from all attributes (i.e. , each query has between 1 and 3 required features on 1, 2, 3 or 4 attributes).

#### Quantitative Analysis

The CIC dataset is summarized by the SAINTETIQ summarization process as described in Section 3. The whole summarization is completed in 9min. The dataset, consisting of 33735 records, yields a summary tree with 13263 nodes, 6701 leaves, maximum depth of 16, average depth of 10.177, maximum width of 14 and an average width of 2.921. The data distribution in the summary tree reveals a 0.6% ( $\frac{6701}{1036800}$ ) occupation rate.

From the analysis of theoretical complexities, we claim that ESA and ESRA algorithms are much faster than the post-clustering process ESA+SEQ. That is the main result of Figure 6 that shows the performance evolution according to the number of cells populated by query answer records. Furthermore, we plot in this figure the number of tree nodes visited (#Visited Nodes) per query (right scale) and finally the normalized ESRA time cost ( $t_{N.ESRA}$ ) to evaluate the performance of the ESRA regardless of how query fits the pre-clustering.  $t_{N.ESRA}$  is computed as follows:

$$t_{N.ESRA} = \frac{t_{ESRA} \cdot \#Tree\ Nodes}{\#Visited\ Nodes}.$$

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<sup>1</sup><http://trec.nist.gov>

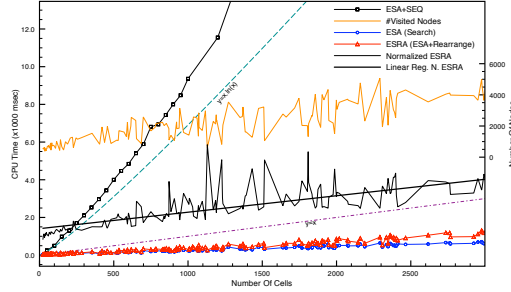


Figure 6: Time cost comparison

As one can observe, the ESA+SEQ approach is quasi-linear ( $O(L \cdot \log(L))$ ) in number of cells  $L$  whereas ESA, ESRA and N. ESRA processes are linear ( $O(L)$ ). Besides, the time cost incurred by rearranging query results (i.e.,  $t_{\text{ESRA}} - t_{\text{ESA}}$ ) is insignificant compared to search cost (i.e.,  $t_{\text{ESA}}$ ). Thus, the ESRA algorithm is able to drastically reduce time cost of clustering query results.

### Qualitative Analysis

Due to the difficulty of conducting a large-scale real-life user study, we discuss the effectiveness of our clustering process of query answers (ESRA) based on structural properties of results provided to the end-user. It is worth noticing that the end-user benefit is proportional to the number of items (clusters or tuples) the user needs to examine at any time as well as dissimilarity between these items.

We define the ‘Compression Rate’ ( $CR$ ) as the ratio of the number of clusters returned as a starting point for an online exploration over the total number of cells covered by such clusters. As expected, Figure 7 shows that  $CR$  values of ESRA and ESA+SEQ approaches are quite similar and much smaller than that of ESA process. Note that  $CR = 1$  means no compression at all, whereas smaller values represent higher compression. As one can observe, size reduction is clearly achieved by the ESRA process.

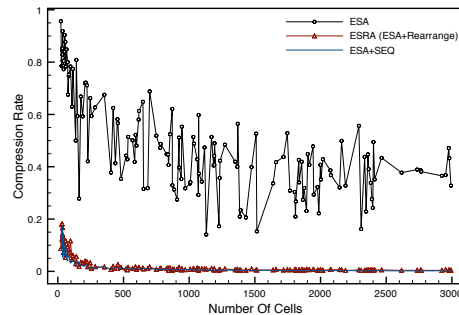


Figure 7: Compression rate comparison



We could also see that the dissimilarity (Figure 8) of the first partitioning that ESRA process presents to the end-user is greater than that of ESA approach and is in the same order of magnitude than that provided by the ESA+SEQ approach. It means that ESRA process significantly improves discrimination between items when compared against ESA approach and is as effective as the post-clustering approach ESA+SEQ. Furthermore, the dissimilarity of the ESA result is quite similar to that of the most specialized partitioning (set of cells). Thus the rearranging process is highly required to provide the end-user with well-founded clusters.

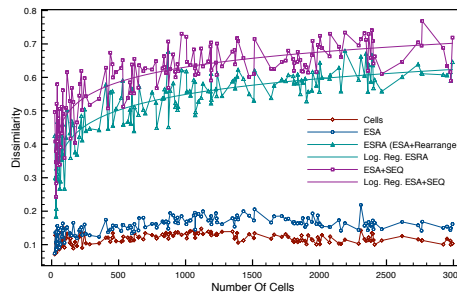


Figure 8: Dissimilarity comparison

In the following, suppose a user considers that a summary  $z$  seems relevant (exactly or partially) to her. She explores  $z$  using either SHOWTUPLES or SHOWCAT option until she reaches every relevant tuple. Thus, the number of hops  $NoH$  (i.e., the number of SHOWTUPLES/SHOWCAT operations) required to reach relevant data ranges from 1 up to  $d_z + 1$ , where  $d_z$  is the height of the tree rooted by  $z$  (i.e., subtree  $H_z$ ). The best case occurs when  $z$  exactly meets user's need whereas the worst case occurs when the relevant information is reached by following a path of maximal length in  $H_z$ . Note that these two scenarios are on opposite extremes of the spectrum of possible situations: the generalized case is that the user's requirement is exactly satisfied by a node (i.e., not necessarily the root or a leaf node) of  $H_z$ . In this experiment, one considers a user query  $q$  with selectivity of  $\eta$  percent, i.e., the ratio of the number of cells covered by  $q$  results over the number of cells of the pre-computed tree. We look for all the possible summaries (subtrees) in the pre-computed tree that can be returned as the result of  $q$  and, for each one, we compute its maximum depth, its average depth and the average length of all paths emanating from that node (summary). Then, we pick out the highest (maximum) value observed for each of these measures. The three values obtained, for each value of  $\eta$  ( $\eta \in \{0.2, 0.4 \dots 2\}$ ), evaluate respectively: (1) the worst number of hops required to reach the deepest leaf (cell) node (Worst NoH2C) containing relevant data; (2) the average number of hops needed to reach any leaf node (Average NoH2C) containing relevant data (i.e. not necessarily the deepest one); (3) the average number of hops required to reach any node (Average NoH2N) containing relevant data (i.e., not necessarily a leaf node).

Figure 9 shows that the Worst NoH2C and the Average NoH2C are relatively high, but bounded respectively by the maximum (16) and the average (10.177) depth of the pre-computed tree. It is worth noticing that, in real life situation,

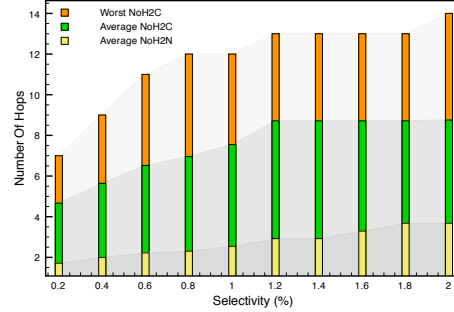


Figure 9: The number of SHOWTUPLES/SHOWCAT operations

the user finds out that her need has successfully served by an inner node of the tree rooted by  $z$ . Thus, the Average NoH2N is more adapted to evaluate the effectiveness of our approach (ESRA). As one can observe, the Average NoH2N is quite small (e.g. , 3.68 hops to locate relevant information within a collection of 674 records).

Finally, we aim to compare the number of results the user has to examine until she reaches relevant data when answers are rearranged using ESRA or not. To this end, we pick randomly one summary  $z$  from the lower part of the pre-computed tree (e.g. ,  $z$  covers 30 cells) and generate the query  $q$  that returns  $z$ . Assume that  $z$  exactly fits user's need, i.e. , the user is interested in all tuples ( $R_z$ ) that mach  $q$  ( $Precision = Recall = 1$ ). Further, assume that such user submits a broad query  $q'$  instead of  $q$ , i.e. ,  $q'$  is a relaxation of  $q$  and consequently matches a superset of  $R_z$  ( $Precision < 1$  and  $Recall = 1$ ). In this experiment, the additional selection predicates in the query  $q$  are used to model user's actions to get exactly what she wants when browsing through  $q'$  results. Note that when no processing has been performed on  $q'$  results, the user has to examine  $N$  cells in order to get all relevant tuples ( $R_z$ ), where  $N$  is the number of cells that satisfy  $q'$ . However, when  $q'$  is processed using our ESRA algorithm, the user has to examine only  $ENoH$  clusters and then performs one SHOWTUPLES operation;  $ENoH$  is the effective number of hops (or SHOWCAT operations) required to reach  $z$ . This is due to the fact that our ESRA algorithm provides a clustering schema of  $q'$  results where all cells that fit user's need are grouped into a single cluster (i.e.  $z$ ) which does not overlap with irrelevant ones, if we except overlapping of borders of fuzzy cells. Figure 10 shows the evolution of  $\frac{ENoH}{N}$  according to the precision of  $q'$ . The precision of  $q'$  is the ratio of the number of relevant cells (30) over the number of cells that match the broad query  $q'$  ( $N$ ). As one can observe,  $\frac{ENoH}{N}$  is much smaller than 1. For instance,  $\frac{ENoH}{N}$  is equal to 0.0013 ( $\frac{5}{3603}$ ) for a query precision of 0.0083 ( $\frac{30}{3603}$ ). It means that only 5 hops are required to locate the summary  $z$  within a clustering schema of  $q'$  results that covers 3603 cells. Thus, ESRA is able to drastically reduce the 'Information Overhead' that user experiences when submitting a broad query.

Those experimental results validate the claim of this work that is to say the ESRA algorithm is very efficient and achieves high quality results and consequently makes the exploration process more effective.

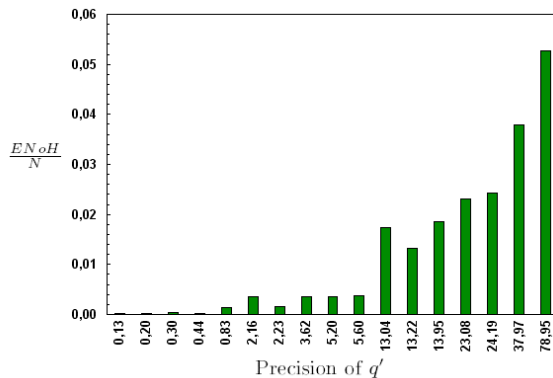


Figure 10: Effectiveness of ESRA regarding the precision of query's results

## 8 Conclusion

Interactive and exploratory data retrieval are more and more suitable to database systems. Indeed, regular ‘blind’ queries often retrieve too many answers. Users then need to spend time sifting and sorting through this information to find relevant data. In this communication, we propose an efficient algorithm coined Explore-Select-Rearrange Algorithm that quickly provide the user with concise and well-formed clustering schema of a query results. Indeed, clusters of a user query results are targeted elements of the hierarchical grid-based clustering of the whole data set (computed offline) and every subtree rooted by each answer offers a guided tour of the data subset to the user. Experimental results show that our process is efficient, provides well-formed clusters (their number is small and they are well separated) and then is very helpful in a situation when a user has not a specific retrieval goal and is interested in browsing through a set of items to explore what choices are available.

Note that Explore-Select-Rearrange Algorithm, in its current state, returns results using linguistic variables of summaries even if user expresses her query using a free vocabulary. A future work is then to perform results rewriting.

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## Contents

<b>1</b>	<b>Introduction</b>	<b>3</b>
<b>2</b>	<b>Related work</b>	<b>4</b>
<b>3</b>	<b>Overview of the SAINTETIQ System</b>	<b>5</b>
3.1	A Two-Step Process . . . . .	5
3.1.1	Mapping Service . . . . .	5
3.1.2	Summarization Service . . . . .	6
3.2	Features of Summaries . . . . .	7
3.3	Discussion about SAINTETIQ . . . . .	8
<b>4</b>	<b>Querying the SAINTETIQ Summaries</b>	<b>8</b>
4.1	Expression and Evaluation of a Query . . . . .	10
4.2	Search algorithm . . . . .	10
4.3	Multi-Scale Summarized Answers . . . . .	12
<b>5</b>	<b>Rearranging query results</b>	<b>14</b>
<b>6</b>	<b>Discussion about Extension to any Fuzzy Predicate</b>	<b>16</b>
6.1	Query rewriting . . . . .	17
6.2	Sorting the results . . . . .	19
<b>7</b>	<b>Experimental Results</b>	<b>20</b>
7.1	Data Set . . . . .	20
7.2	Results . . . . .	20
<b>8</b>	<b>Conclusion</b>	<b>24</b>



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