HASAR: Mining Sequential Association Rules for Atherosclerosis Risk Factor Analysis

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Abstract. We present the HASAR method that is an hybrid approach for extracting adaptive sequential association rules. This method extracts association rules between events occurring in subsequent time-intervals using closed itemsets extraction and evolutionary techniques. An important feature is its capacity to consider different time-intervals depending on the attributes semantic. We applied this method for the analysis of long term medical observations of atherosclerosis risk factors for cardio-vascular diseases prevention. Experimental results show that it is well-suited for extracting knowledge from temporal data where interesting patterns have different observation period length.

1 Introduction

In this paper, we consider sequential association rules which express casualty relationships between sets of events occurring in subsequent time-intervals. We developed a specific approach, called HASAR for Hybrid Adaptive Sequential Association Rules, for finding such patterns of interest. Our methodology is applied to an health care problem, but it is also suited to a broad collection of data mining problems where data are temporal observations on individuals, in customer behaviour or credit risk prediction for instance. Many studies have focused on efficient mining of sequential patterns or patterns in time-related data. Most of them are based on extensions of the APRIORI algorithm [AMS96] proposed for extracting association rules. The HASAR approach presented in this paper combines techniques for searching closed itemsets, as defined in the CLOSE algorithm [PTB04], and an heuristic approach relying on a genetic algorithm.

We used the HASAR approach for the analysis of long term observation data in the STU-LONG dataset. This dataset was constituted in the framework of a longitudinal study of atherosclerosis risk factors to evaluate the impact of non-pharmacological prescriptions on these risks. It contains data collected between 1975 and 2001 on a population of 1417 men born between 1926 and 1937 in Czechoslovakia. First, an entry examination was performed and data concerning social characteristics, diet, tobacco and alcohol consumption, physical activities, personal anamnesis and, physical and biochemical examinations were collected. During this examination, patients were classified into three groups according to principal atherosclerosis risk factors (RFs): Arterial hypertension, hypercholesterolemia, hypertriglyceridemia, overweight, smoking and positive family case history. These three groups are the following:

– Normal group (NG): No risk factor and no cardio-vascular disease.
– Pathological group (PG): Cardio-vascular disease diagnosed.

During the twenty one years following the patients’ entry, control examinations were performed to record changes in diet, smoking habits, physical activities and responsibility level in job, sport practice in leisure time and, physical and biochemical measures. For each control, the patient id, the date and the control number were recorded. In 2001, 389 patients were deceased and, the cause and date of death were recorded. For a group of 403 patients that answered to a postal questionnaire, detailed data – similar to those gathered during controls – were collected.

The STULONG dataset was collected at the 2nd Department of Medicine, 1st Faculty of Medicine of Charles University and Charles University Hospital, U nemocnice 2, Prague 2 (head. Prof. M. Aschermann, MD, SDr, FESC), under the supervision of Prof. F. Boudík, MD, ScD, with collaboration of M. Tomečková, MD, PhD and Ass. Prof. J. Bultas, MD, PhD. The data were transferred to the electronic form by the European Centre of Medical Informatics, Statistics and Epidemiology of Charles University and Academy of Sciences (head. Prof. RNDr. J. Zvárová, DrSc.). At present time the data analysis is supported by the grant of the Ministry of Education CR Nr LN 00B 107.

Objectives. One of the main objectives of this study was to evaluate the impact of behavioural changes – starting or stopping a diet or sport practice for instance – on the RF and the development of cardio-vascular diseases (CVDs). The initial analytical questions in STULONG evolved after the first experimentations and discussions with medical experts, mainly because of the evolutions of medical knowledge since the beginning of the study and missing data (e.g., uric acid measure is given for less than 10% of controls).

During this work, we have developed a new method that we believe will help to answer the following questions defined throughout discussions with medical experts and related to the long-term observations:

– Are there differences between men of the normal, risk and pathological groups from the viewpoint of the impact of behavioural changes on RF and CVD development?
– What characterizes men who developed a CVD and those who stayed healthy on the global population and the risk group?
– Are the education level and the responsibility in job good criteria for segmenting patients with perilous or safe behaviours and high or low RF?

HASAR is an association rules extraction approach incorporating temporal relationships. It extracts sequential association rules which are well fitted to analyse casualty relationships between behavioural changes and RF evolutions or CVD development. Association rule extraction, first introduced in [AIS93], aims at discovering casualty relationships between sets of attribute values, called itemsets, in large datasets. An example association rule, fitting in the context of market basket analysis is:

\[ \text{BUY(cereal)} \land \text{BUY(sugar)} \rightarrow \text{BUY(milk)}, \text{support} = 20\%, \text{confidence} = 75\%. \]

This rule states that customers who buy cereal and sugar also tend to by milk. The support measure indicates that 20% of all customers bought both three items and the confidence measure shows that 75% of customers who bought cereal and sugar also bought milk. Informally, the support represents the range of the rule and the confidence indicates the precision of the rule. In order to extract only statistically significant association rules, only those with support and confidence at least equal to some user defined minsupport and minconfidence thresholds are generated.
Organization. In section 2, we present preparation and transformation methods applied to the dataset for extracting long-term observation related patterns. Sequential association rules and the techniques we used for their extraction are defined in section 3. In section 4, we show experimental results and section 5 concludes the paper.

2 Data preparation

Data for the long-term observation were collected during the twenty one years controls after the patients' entry. We used both entry and control data to generate multiple datasets that are adapted to the kind of knowledge we were interested in. Attributes of interest were selected and prepared according to discussions with medical experts, for determining threshold values of physical and biological measures for instance.

2.1 Sequential rules and search strategy

Since our main objective concerns the effect of behaviour on risk factor development, we decided to look for sequential rules involving casualty relationships between patient behavioral changes and risk factor changes on subsequent time intervals. Sequential rules with the form \( X \rightarrow Y \) we search for involve both itemsets and time-itemsets. We call \( \text{time\_item} \) an attribute value occurring in a particular temporal window. We call \( \text{time\_itemset} \) a set of \( \text{time\_items} \). Our sequential rules have the following structure:

\[
\text{IDE\_itemset} \land \text{BEH\_time\_itemset} \rightarrow \text{RF\_time\_itemset}
\]

where the components are:

- \( \text{IDE\_itemset} \): an itemset of static identification attributes,
- \( \text{BEH\_time\_itemset} \): a time itemset of behavioural attributes,
- \( \text{RF\_time\_item} \): a time item on a risk factor attribute.

An example sequential rule may be:

\( \text{ALCOHOL=regularly} \land \text{BEH\_PHA=decreased\_sits} \rightarrow \text{RF\_CHOLEST=increased} \).

Such a rule must be interpreted as a casualty relationship between changes on risk factors occurring on an \( \text{observation\ temporal\ window} \) of \( O \) months and induced by static data and by changes on behaviour on a previous \( \text{action\ temporal\ window} \) of \( A \) months. We also defined a \( \text{latency\ temporal\ window} \) \( L \) between the \( \text{action\ period} \) and the \( \text{observation\ period} \) which allows a waiting time to observe the impact of some behavioural changes.

The example rule should be interpreted as follows:

\[
\text{if the patient regularly consumed alcohol when he entered the study and his physical activity after job decreased over a } A \text{ month period then his cholesterol rate increased at a control which occurred } L \text{ months after and over the subsequent } O \text{ month observation time.}
\]

An important element for the method flexibility is that temporal window sizes \( O, A \) and \( L \) are defined as parameters of sequential rules. The strategy we applied for extracting these rules was first running wide data transformations to tailor data to the specific task. This first step consisted in applying corrections, replacing missing values, creating new attributes and a new table for saving behavioural and risk factor changes and flattening data on changes.
Rule quality is computed according statistical criteria like traditional association rules. Statistical measures are presented in sections 3.1 and 3.3. HASAR flexibility is also provided by the evolutionary algorithm involved for searching for rules. We defined a Genetic Algorithm (GA) for this task. GAs [GR87] are well suited to large combinatorial search spaces. GAs are adaptive procedures that evolve a population of structures in order to find the best individual. The evolution is performed by specific genetic operators like mutation and crossover. They have a long history of being exploited for rule manipulation [Freitas02,SAL03]. As it was suggested, they offer techniques such as niching which allow not only to find the best rule, but a selection of good rules.

2.2 Patient classes

In order to observe potential differences between classes of patients, we distinguished the following partitions:

- Groups NG, PG and RG assigned to patients in the STULONG study.
- Classes CVD and NCVD which respectively represent patients who had and did not have a cardio-vascular disease during the STULONG study.
- Clusters of patients based on their education level and job responsibility criteria.

Classes CVD and NCVD were obtained by splitting the EXPERIMENT table using attributes CONTROL.H0Dnx, with $x \neq 0$ and $x \neq 15$, that indicates if a cardio-vascular disease was diagnosed and DEATH.PRICUMR that indicates if his death was due to a cardio-vascular disease.

Clusters of patients based on their education level and job responsibilities criteria were obtained as follows. In a first attempt, we extracted all closed patterns containing at least the social factors but those gathering a relevant number of patients (at least 200) did not reveal a significant medical interpretation. This is due to the fact that some attribute values cover a large number of patients (e.g., 1023 patients among the 1199 are married).

After talking with a physician, the ‘education level’ (VZDELANI) and “responsibilities in job” (ZDP0V) attributes, that are most likely to influence atherosclerosis, were chosen as main criteria for building the clusters. We investigated the closed patterns containing the items coming from these attributes. We got the following 11 closed patterns (or potential clusters):

1. basic school and others (for responsibilities in job)
2. primary school and managerial worker
3. primary school and partly independent worker
4. primary school and others
5. secondary school and managerial worker
6. secondary school and partly independent worker
7. secondary school and others
8. university and managerial worker
9. university and partly independent worker
10. university and others
11. university and pensioner (not because of ICHS)

The closed patterns number 5 and 8 were merged to produce the first cluster. We perform a similar process with closed patterns number 6 and 9, 7 and 10, 1 and 4. The fifth cluster contains the 151 remaining transactions (closed patterns 2, 3 and 11). Finally, we obtain the non-overlapping clusters described in table 1.
<table>
<thead>
<tr>
<th>Cluster</th>
<th>Secondary school</th>
<th>Responsibility in job</th>
<th>Number of patients</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Healthy</td>
</tr>
<tr>
<td>1</td>
<td>yes</td>
<td>managerial worker</td>
<td>150</td>
</tr>
<tr>
<td>2</td>
<td>yes</td>
<td>partly independent worker</td>
<td>227</td>
</tr>
<tr>
<td>3</td>
<td>yes</td>
<td>others</td>
<td>127</td>
</tr>
<tr>
<td>4</td>
<td>no</td>
<td>others</td>
<td>221</td>
</tr>
<tr>
<td>5</td>
<td>-</td>
<td>-</td>
<td>94</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Total of patients</td>
<td>819</td>
</tr>
</tbody>
</table>

Table 1. Description of clusters.

2.3 Attributes of change and new tables

A preliminary step consisted in correcting some errors or contradictions and replacing missing and "not stated" values by the same value. We built new tables CHANGES and EXPERIMENT more fitted to the task from the initial tables ENTRY, CONTROL. The DEATH table was used in order to split the patient set into the CVD and NCVD classes. Discussions with medical experts allowed us to identify some guidelines for building tables and to understand which initial attributes we had to keep and which ones we have to build. First, we kept existing identification attributes (IDE attributes) about patients. These attributes are named according to the expression `IDE ATTRIBUTE NAME`. They represent:

- the education level of the patient,
- the age of the patient at the moment of the control,
- the initial group of the patient when he came into the study,
- the alcohol consumption at the beginning of the study since STULONG data do not provide this information for each control.

Other attributes describe behavioural changes and changes related to risk factors from one control to another. Behavioural change attributes (BEH change attributes) were named according to the expression `BEH ATTRIBUTE NAME` and risk factor change attributes (RF change attributes) were named according to the expression `RF ATTRIBUTE NAME`. Attributes for behavioural changes are related to criteria below:

- consumption of cigarettes a day,
- physical activity in job and after job,
- different kinds of diet,
- medicine for cholesterol and for blood pressure.

Attributes for risk factor changes are related to the following criteria:

- global, HDL and LDL cholesterol levels,
- triglycerides level,
- overweight or obesity,
- blood pressure measures,
- glycemia level.

The new CHANGES table is composed with IDE, BEH change and RF change attributes. For two subsequent controls number N and N+1 of a patient, a new tuple is created and the value of each BEH and RF attribute is computed by comparing attribute values of the two controls. For the first control, a new tuple is created by comparing it with information in the ENTRY table for the patient. CHANGES thus contains as many tuples as the CONTROL table. As an example, we show in table 2 and table 3 how the BEH_PHA and
<table>
<thead>
<tr>
<th>Value</th>
<th>CONTROL.AKPOZAM [N]</th>
<th>CONTROL.AKPOZAM (N+1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>stay_sits</td>
<td>he mainly sits</td>
<td>he mainly sits</td>
</tr>
<tr>
<td>decreased_sits</td>
<td>moderate or great activity</td>
<td>he mainly sits</td>
</tr>
<tr>
<td>increased_modest</td>
<td>he mainly sits</td>
<td>moderate activity</td>
</tr>
<tr>
<td>stay_modest</td>
<td>moderate activity</td>
<td>moderate activity</td>
</tr>
<tr>
<td>decreased_modest</td>
<td>great activity</td>
<td>moderate activity</td>
</tr>
<tr>
<td>increased_great</td>
<td>he mainly sits or moderate activity</td>
<td>great activity</td>
</tr>
<tr>
<td>stay_great</td>
<td>great activity</td>
<td>great activity</td>
</tr>
</tbody>
</table>

Table 2. Attribute BEH.PHA.

<table>
<thead>
<tr>
<th>Value</th>
<th>CONTROL.CHLST (N)</th>
<th>CONTROL.CHLST (N+1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>stay_normal</td>
<td>&lt; 6</td>
<td>&lt; 6</td>
</tr>
<tr>
<td>decreased</td>
<td>≥ 6</td>
<td>&lt; 6</td>
</tr>
<tr>
<td>increased</td>
<td>&lt; 6</td>
<td>≥ 6</td>
</tr>
<tr>
<td>stay_high</td>
<td>≥ 6</td>
<td>≥ 6</td>
</tr>
</tbody>
</table>

Table 3. Attribute RF.cholesterol.

RF.cholesterol attribute values are computed from values of attributes CONTROL.AKPOZAM (which indicates the physical activity after job) and CONTROL.CHLST (which indicates the global cholesterol rate).

In order to get a temporal description of each patient all along the study, we flattened the CHANGES table. The EXPERIMENT table is the result of the flattening operation; it contains as many tuples as patients in CONTROL. One tuple in EXPERIMENT contains IDE attributes of the patient and changes attributes for every control the patient passed through.

3 Strategies

3.1 Target model and definitions

We consider the time segment which represents the intervention on a patient from the time he entered the study until the time he left it. Our discussions with medical expert led us to fix a month as the time unit. Let us consider a patient and the time-interval [in; out] during he was in the STULONG study.

\( C(T, \text{patient}) \) refers to a control occurring at time \( T \) months after \( in \) for the patient. A control \( C(T, \text{patient}) \) is characterized by a \( \text{BEH\_time\_itemset} \) and a \( \text{RF\_time\_item} \) which represents behavioural and risk factors changes observed for the patient at this control time. A control period \( CP(T, \text{patient}) \) is a \( A \)-month size time interval \([T; T+A]\) where it occurs one control at least for the patient. A temporal configuration \( TC(T_1, T_2, \text{patient}) \) is a time interval \([T_1; T_2]\) such as:

- it exists a control period \( \gamma = CP(T, \text{patient}) \) with \( T_1 \in [T; T+A] \),
- \( T_2 \in [T_1 + L; T_1 + L + O] \),
- it occurred a control \( C(T_2, \text{patient}) \),
- it did not occur any control in the interval \([T + A; T_2]\) for the patient.

We say that two temporal configurations \( TC(T_1, T_2, \text{patient}) \) and \( TC(T_3, T_4, \text{patient}) \) are compatibles if \( T_2 \leq T_3 \) or \( T_4 \leq T_1 \).

**Statistical measures.** We define the measure \( \text{max\_support} \) as the whole number of possible compatible temporal configurations for all records (patients) in the flattened table EXPERIMENT. For a rule antecedent \( X = IDE\_itemset \cup BEH\_time\_itemset \), we say that a temporal configuration \( TC(T_1, T_2, p) \) contains \( X \) if:
- IDE\textunderscore itemset is observed for the patient $p$.
- BEH\textunderscore time\textunderscore itemset is observed over a control period $\gamma = CP(T, patient)$.

For a rule consequent $Y = RF\textunderscore time\textunderscore item$, we say that a temporal configuration $TC(T_1, T_2, p)$ contains $Y$ if $RF\textunderscore time\textunderscore item$ is observed at control $C(T_2, patient)$. For a rule antecedent $X$, we define the cardinality measure of $X$, $\text{card}(X)$, as the number of possible compatible $TC(T_1, T_2, p)$ that contains $X$. For a rule consequent $Y$, we define the cardinality measure of $Y$, $\text{card}(Y)$, as the number of possible as the number of possible compatible $TC(T_1, T_2, p)$ which contains $Y$. For a rule $X \rightarrow Y$, we define the cardinality measure of $X \land Y$, $\text{card}(X \cup Y)$ as the number of possible $TC(T_1, T_2, p)$ that contains $X$ and $Y$.

### 3.2 Association rules based approach

Two main approaches for extracting association rules can be distinguished. In the first approach, all itemsets with $\text{support} \geq \text{minsupport}$, called frequent itemsets, are extracted and all association rules with $\text{confidence} \geq \text{minconfidence}$ are generated from them. This approach is very efficient when data are weakly correlated, such as market basket data, but performances drastically decrease when data are dense or correlated, such as statistical data for instance. A comprehensive survey of this approach can be found in [AMS+96].

The second approach is based on the extraction of generators and frequent closed itemsets defined using the Galois closure operator. From these, the informative basis for association rules containing non-redundant association rules with minimal antecedent and maximal consequent. This approach both improves the extraction efficiency, by reducing the search-space, and the result relevance, by suppressing redundant rules, in the case of dense or correlated data. A summary of this approach can be found in [PTB+04].

**Frequent closed itemsets and generators.** Frequent closed itemsets and generators are defined according to the closure operator $\phi$ of the Galois connection. This operator associates with an itemset $l$ its closure $\phi(l)$ that is the maximal set of items common to all objects containing $l$. That is, the closure of $l$ is the intersection of all objects containing $l$. The minimal closed itemset containing an itemset $l$ is its closure $\phi(l)$ and we say that an itemset $l$ is a closed itemset if $\phi(l) = l$. The generators of a closed itemset $c$ are the minimal\footnote{With respect to the inclusion relation.} itemsets which closure is $c$. Generators are the minimal itemsets we can consider for discovering frequent closed itemsets, by computing their closures. Since the support of a frequent itemset is equal to its closure support and since maximal frequent itemsets are maximal frequent closed itemsets, the frequent closed itemsets constitute a minimal non-redundant generating set for all frequent itemsets and thus, for all association rules. Consider the dataset $D$, constituted of six objects identified by their OID and five items, represented in figure 1(a). The eight generators and five frequent closed itemsets, with their supports, in $D$ for $\text{minsupport} = 2/6$ are given in figure 1(b).

The itemset $\{A\}$ is the generator of the frequent closed itemsets $\{AC\}$: the intersection of all objects containing $\{A\}$, that are objects 1, 3 and 4, gives $\{AC\}$ and no subset of $\{A\}$ has $\{AC\}$ as closure. $\{A\}$ and $\{AC\}$ both have a support of $\frac{3}{6} = 1/2$. The frequent closed itemset $\{BCE\}$ has two generators: $\{BC\}$ and $\{CE\}$. $\{BE\}$ is not a generator of $\{BCE\}$ since it is a frequent closed itemset, $\{B\}$ and $\{E\}$ are generators of $\{BE\}$ and $\{C\}$ is itself its own generator.
3.3 Evolutionary approach

Genetic Algorithms are robust, flexible algorithms which tend to cope well with attribute interaction in atherosclerosis data. Furthermore, the comprehensibility of the discovered knowledge is important and GAs allow us to extract comprehensible rules evolving populations of prediction patterns. Our GA implementation uses EO, a templates-based, ANSI-C++ compliant evolutionary computation library.

**Genome.** The first issue in designing a GA is how to encode each individual in the population. To represent variable length rule we use a fixed-length genome which contains a gene for each IDE _attribute_ and BEH _attribute_ and another gene for one of the RF _attributes_. Each gene contains three elements: attribute name, attribute value and an activation flag indicating whether or not an item in the rule is associated to the gene.

**Generation of initial population.** The method used to generate the initial population is based on CLOSE algorithm results. CLOSE generate a list with generators and their frequent closed itemset associated, that allows us to initialize the population in three steps:

- Rule antecedents are created from generators. Generators containing only RFs are skipped.
- Rule consequents are created with the first RF found in the frequent closed itemset or the generator. If no RF is found, we generate randomly one.
- For each attribute not represented in generators the value of the corresponding gene is randomly defined and the activation flag is set to false.

**Genetic operators.** We use a tournament selection with size of 2. The selection is deterministic, starting from the best ones down to the worse ones. If the total number to select is less than the size of the source populations, the best individuals are selected once. If more individuals are needed after reaching the bottom of the population, then the selection starts again at top. If the total number required is N times that of the source size, all individuals are selected exactly N times. For replacement we use the most straightforward method, called generational replacement where all offspring replace all parents; however weak elitism is used.

New patterns are generated by combining existing patterns using a crossover operator or by modifying existing patterns via a mutation operator. Crossover is a recombination operator that swaps genetic material between two individuals. We used a one point crossover method. Three mutation operators were used: first one simple changes the
attribute of a gene with a random attribute, the second generates a random transition value (domain of this value is a parameter of the GA) which is randomly added or subtracted to the current gene value and the last inverts the current value of the activation flag. All of the mutation rates can be define independently.

**Fitness function.** A crucial issue in the design of a GA is the choice of the fitness function. In a first approach we only consider support to select the most frequent rules, confidence to consider reliable rules and lift to ensure a high level of dependence between antecedent and consequent part of a rule. To evaluate the quality of a rule $r: X \rightarrow Y$, our GA applies the fitness function on the individual associated to the rule.

$$fitness(r) = support(r) \times confidence(r) \times lift(r)$$  \hspace{1cm} (1)

$$support(r) = \frac{\text{card}(X \cup Y)}{\text{max}_r \text{support}}$$ \hspace{1cm} (2)

$$confidence(r) = \frac{\text{card}(X \cup Y)}{\text{card}(X)}$$ \hspace{1cm} (3)

$$lift(r) = \frac{\text{card}(X \cup Y)}{\text{card}(X) \times \text{card}(Y)}$$ \hspace{1cm} (4)

4 Experimental results

4.1 Patients classes comparison

**Patient groups.** The experience consisted in extracting rules on PG and testing them on NG and RG. Differences for support and fitness measures are shown in figure 2. One may observe that the best rules found on PG are not valid on NG. Most of them have a good support but a weak fitness on RG. Thus, these results show quite different relationships between the patient behaviour and their risk factors among initial groups.

![Graph showing support and fitness](image)

**Fig. 2.** Best rules on PG versus NG and RG.

**Cardiovascular disease.** The experience consisted in extracting rules on CVD and testing them on NCVD. Results in figure 3 show that the best rules found on CVD have similar support and fitness on NCVD. These results seem to show that relationships between behavioural changes and risk factors are not really different between CVD and NCVD patients.
Fig. 3. Best rules on CVD versus NCVD.

Social clusters. This experience consisted in extracting rules on Cluster1 and testing them on Cluster3 and Cluster4. We can see in figure 4 that the best rules found on Cluster1 also have good measures on Cluster4 and results on Cluster3 are quite similar. Thus it seems that social factors like education level and job responsibility do not allow to distinguish different behaviour from the viewpoint of cardiovascular risks.

Fig. 4. Best rules on Cluster1 versus Cluster3 and Cluster4.

4.2 Initialisation methods

In order to evaluate performances of our initialization method we compare it with a random initialization method. Results for PG and RG groups are shown in figure 5 and results for CVD and NCVD are shown in figure 6. The first three columns show statistics about initial populations and the last column show the fitness of best individuals after a GA run. We can observe that mean fitness of populations generated using CLOSE is 8.75 to 400 times better than those of randomly generated population. It is not surprising since CLOSE optimize rules support and confidence, two means criteria of our fitness function. However, it is interesting to note that after a GA run the fitness of the best offspring of populations generated using CLOSE is 1.55 to 4.79 times better than the best offspring of randomly generated population. Furthermore, an analyze of rules show that GA doesn’t converge toward local optima given by CLOSE. Then we succeed in improving GA performances conserving diversity in solutions proposed.
Fig. 5. Comparing initialization methods on patient groups.

Fig. 6. Comparing initialization methods on patient classes.

4.3 Observation time windows variations

Time windows $A$, $L$ and $O$, used to extract antecedents and consequents of rules, define time itemsets. An important aspect of the HASAR method is its capacity to make vary these time windows. In the STULONG analysis, the observation time of RF evolutions depends on the considered risk. For instance, according to physicians' knowledge, the effects of a diet on the weight are perceptible after a few months whereas the effects on the cholesterol measures are most often perceptible after a longer period.

We evaluated the effect of RFs observation time window variations on the fitness of rules for two RFs: RF_CHOLES and RF_BLOODPRESS. The results are shown in figure 7. For RF_CHOLES, rules generated for a 60 months window have a much better fitness than those generated for a 15 months window. For RF_BLOODPRESS, the situation is the opposite: a 15 months window gives better fitness than a 60 months window. This shows that effects of non pharmacological prescriptions on hypercholesterolemia must be observed on much longer time period than effects on arterial hypertension.

5 Conclusion

In this paper, we have presented an innovative method for extracting adaptive sequential rules. We have applied the method on the atherosclerosis STULONG dataset. While
previous works on this dataset essentially focused on static information from the ENTRY
Table, we have investigated the analysis of temporal data in the CONTROL table. Our
approach is based on two main points:

- a set of data transformations which is suited to various temporal data,
- an hybrid strategy which combines advantages from quite different techniques: an
  exhaustive search for frequent generators and closed itemsets which are used as the
  initial population of an evolutionary algorithm.

Experimental results allowed to point out different tendencies among patient groups
and confirmed prior medical knowledge. In order to answer in-depth to the analytical
questions, further investigations of sequential rules with the assistance of medical experts
are required.

In the future, we plan to apply the HASAR approach to other temporal datasets where
observation time windows are not uniform on all attributes. Another interesting perspective
is to extend the approach by integrating background knowledge, such as expressed
in medical ontologies, in the search process.

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