



# Gastric Cancer Risk Analysis in Unhealthy Habits Data with Classification Algorithms

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Abstract – Data mining methods are applied to a medical task that seeks for the information about the influence of Helicobacter Pylori on the gastric cancer risk increase by analysing the adverse factors of individual lifestyle. In the process of data preprocessing, the data are cleared of noise and other factors, reduced in dimensionality, as well as transformed for the task and cleared of non-informative attributes. Data classification using C4.5, CN2 and k-nearest neighbour algorithms is carried out to find relationships between the analysed attributes and the descriptive class attribute – Helicobacter Pylori presence that could have influence on the cancer development risk. Experimental analysis is carried out using the data of the Latvian-based project "Interdisciplinary Research Group for Early Cancer Detection and Cancer Prevention" database.

*Keywords* – Classification, data pre-processing, gastric cancer risk analysis.

#### I. INTRODUCTION

One of the leading causes of gastric cancer is the *Helicobacter Pylori* (*H. pylori*) bacteria. *H. pylori* is bacterium with spiral form that lives in the mucous lining of the stomach. *H. pylori* has coexisted with humans for many centuries and millennia and infection with these bacteria is very common [1]. *The Centre for Disease Control and Prevention* (*CDC*) affirms that around two thirds of the world population are infected with this bacterium type and infection probability in newly developed countries, including Latvia, is higher than in developed countries [2]. The bacteria spread through contaminated food and water, as well as mouth-to-mouth contact.

Most of the people who are infected with these bacteria do not experience any symptoms caused by bacteria although *H. pylori* can cause chronical active and lasting gastritis, inflammation of duodenum or gastric ulcers in adults, as well as in children. *H. pylori* infection is considered one of the primary causes of gastric cancer. Infection with *H. pylori* is mainly influenced by risk factors like diet, age, history of particular diseases in family, smoking and sanitary conditions [3].

An international study carried out in Latvia has gained data about habits of respondents and their *H. pylori* test results [4]. This article analyses the data obtained in this study.

The article is based on research, where data mining methods and algorithms are used to analyse impact of harmful factors on development of *H. pylori*, which causes higher risk of developing gastric cancer. The aim of the research is to use classification in order to determine the set of harmful factors, which has the highest impact on development of *H. pylori*.

This could help in gastric disease prevention – discovery of potential risk groups would help target tests and therapies and decrease the potential group of screening.

### II. RELATED WORKS

Although there are many studies in the field of medicine, which analyse risk factors for developing diseases and infection, they are specifically designed for factor testing: contaminated food and water, poor hygiene, age, gender, high salt consumption; they either focus on single factor analysis or use only descriptive statistical methods or at most a linear regression analysis. This might not be sufficient to understand complex relationships among all factors. The connected medical research includes information about many different sociodemographic, lifestyle and health factors that have the potential to be significant in H. pylori infection development or lack thereof. Therefore, the study uses easily interpretable machine learning techniques that do not impose specific requirements towards data distributions, matching variances, sample sizes and ratios etc. The authors could not locate any previous articles that would implement such techniques in the corresponding scope [5]–[9].

## III. USED METHODS

The analysed data contain descriptive information about the respondent (135 attributes) and result of the helicobacter test (1 attribute), which can be viewed as the dependent attribute – the class. One of the tasks of data mining is to determine relations in a data set between the set of independent attributes and the dependent attribute. This type of relationships can be discovered using classification [5], [11]. Before beginning the classification process, the data should be pre-processed and attribute informativeness should be determined.

## A. Data Pre-processing

The initial data set obtained from a database can hold noisy, conflicting and missing data. Origins of such data can be various, e.g., most often it stems from the human factor – errors of data input operator or impartial information, as well as incompatible technologies (like language specifics). The use of data mining techniques with such data without processing and preparing them can lead to faulty results; therefore, data should be first analysed using pre-processing techniques. It includes several data processing techniques like data cleaning (replacing missing values by manually entering values or using some constant values instead of the missing values), data reduction (discretization of attribute values and

determining of attribute informativeness) and data transformation (normalisation, construction of new attributes and attribute aggregation), which are described in detail in [5]–[12]. Data pre-processing is one of the most resource-consuming processes – it can take up to 80 % time and work of all data analysis process [4].

In the analysis process of the present research, data are normalized using *z*-score normalisation with standard deviation because this method does not require defining minimum and maximum limits for attributes and it is considered to be one of the most popular data normalisation methods in data mining [13]. A normalized value of attribute  $A_i$  can be calculated as follows (1):

$$A_{i}^{'} = \frac{A_{i} - A}{\sqrt{\frac{1}{n}\sum_{i=1}^{n}(A_{i} - A)^{2}}},$$
(1)

where

 $\overline{A}$  – mean arithmetic value of attribute A;

 $A_i - i$ -th value of attribute A;

n – numbers of records in the data set.

Construction of new attributes is used in order to improve the process of data mining. Attribute construction is carried out by value merging using mathematical operations, e.g., attribute *Weight* and *Height* can be used to construct an attribute named *BMI* [14], which represents body mass index and is calculated as follows in (2):

$$BMI = \frac{Weight}{Height^2},$$
 (2)

where

*Weight* – weight of the respondent, kg; *Height* – height of the respondent, m.

## B. Attribute Selection

Attribute selection decreases the size of a data set and, therefore, increases the speed of data processing algorithm and accuracy of its results [5]. Attribute selection means determining a subset of the most informative attributes. Attribute selection consists of four steps: subset selection, subset evaluation, testing of end criterion and analysis of the obtained results that are shown in Fig. 1.

Selection of the most informative attributes is carried out using *CfsSubsetEval* method, which is considered one of the most popular attribute informativeness evaluation methods used in data mining [11]. Method *Cfs* (Correlation-Based Feature Selection) is an attribute evaluation method based on a correlation analysis.

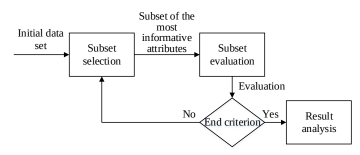


Fig. 1. Selection process of the most informative attributes.

The main idea behind it is to acquire a data set with attributes that have high correlation with class attribute and the smallest possible inter-attribute correlation. The obtained attribute subset is evaluated using (3), which is based on an Equation for Pearson correlation with standardized values [15]:

$$r_{zc} = \frac{kr_{zi}}{\sqrt{k+k(k-1)r_{ii}}},\tag{3}$$

where

 $r_{zc}$  – correlation coefficient (evaluation metric);

k – number of attributes;

 $\overline{r_{ii}}$  – mean correlation of attribute-class pairs;

 $r_{ii}$  – mean correlation of attribute-attribute pairs.

The search algorithm used with *CfsSubsetEval* attribute evaluation method is a *genetic algorithm*. *Genetic algorithm* is a search and optimisation approach, which is based on mechanisms of natural evolution like crossover, mutation and survival of the fittest (more optimal) individuals. Attribute selection using *genetic algorithm* is carried out similarly to a random search. It allows a population, formed by several individuals (solutions), to evolve based on a specific set of rules until a satisfactory result is achieved (being close to optimal result, which can be achieved in affordable time using the available resources). The *genetic algorithm* consists of three steps:

- selection a set of best individuals is selected for a following generation;
- crossover forms new individuals by combining pieces of good individuals that already exist in the population;
- mutation changes individuals by inflicting random changes in order to increase diversity in the population.

Life-cycle of each generation ends by checking the existing population against the end criterion: if the criterion is not met, another population is evolved and changed using the same steps (described previously) and then evaluated. The cycle of the *genetic algorithm* repeats until a pre-defined end criterion is reached [16]. The working principle of the algorithm is depicted in Fig. 2.

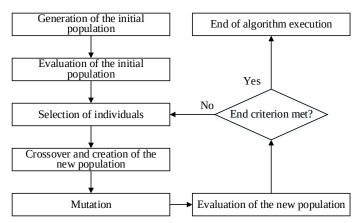


Fig. 2. Steps of the genetic algorithm.

In the beginning, the initial population is generated. It represents a set of possible solutions, where each solution is coded into a chromosome consisting of atomic genes that can be used to form any solution in the possible solution space. The next step involves the evaluation of the population and the use of the best individuals for next generation (the process of selection). Generation is a step in which the same population of individuals is used for operations in order to form new individuals that would be closer to the optimum. The next substeps include crossover and mutation using the individuals of the population, which result in new individuals with different genetic information (different solutions). When each new population is evaluated, it is tested against the end criterion. If the end criterion is met, the algorithm ends its operation. If the end criterion is not met, the algorithm continues with the selection of a new population for next generation [17].

Accuracy of the attribute informativeness evaluation is estimated using *k*-fold cross-validation [11].

## C. Data Classification

Data classification is a process, which produces a model (classifier), which can be used to determine membership of observations to a group or class, by using only a data set with records with determined membership to groups or classes. Classification process consists of two steps:

- Training, where a training data set is used to train a classifier (or build it);
- Testing, where a test data set is used to test the accuracy of the classifier obtained in the first step. If the accuracy of the classifier is sufficient (as determined by system developers or experts), it can be used for classification of new records.

Evaluation of classifier accuracy is carried out using several metrics like *classifier overall accuracy*, *sensitivity* and *specificity*. Values of these metrics determine the suitability of each classifier for solution of the defined task [5], [17].

Simultaneously with the accuracy metric calculations, some approaches for result quality assurance have to be used as well, such as *k-fold cross-validation*. *K-fold cross-validation* divides a data set into *k* subsets by assigning a record to a subset randomly and keeping sizes of subsets equal. Classifier is trained and tested k times. Each time one subset is used for testing and the other k-1 subsets are used for training of a classifier. After each iteration, the results are entered into a confusion matrix, adding them to the results of previous iterations. The final result (confusion matrix and metrics calculated from it) includes results of all k iterations and estimates results of an algorithm as if it were tested on the whole available data set [11].

Experiments with classifiers include applications of the most popular classification algorithms [18]:

- *C4.5*, which uses inductive decision trees as classifiers and is based on algorithm *ID3* with extension that allows it to process data sets that include discrete, continuous and missing data [19];
- *CN2*, which uses inductive reasoning and in each iteration searches for a condition that would cover a large subset of objects with the same class value and as few objects from a different class as possible. When a condition is found, the algorithm removes objects, which are covered by the condition, from the data set and defines a rule that corresponds to the condition [20];
- kNN (*k*-nearest neighbours algorithm), which uses distance metric to determine closeness (distance) between each test object and each training set object and forms a x\*y distance matrix for x objects of the training set and y objects of the test set. For each object from the test set, a set of k neighbours is found by selecting k closest training set objects (based on the smallest distance). The class for each test set record is determined based on a majority vote principle of the neighbour set with or without additional use of heuristics or weighted voting [21].

#### IV. EXPERIMENTS AND RESULT ANALYSIS

Experiments were carried out using data from the database of a study carried out by the Latvian 'Interdisciplinary Research Group for Early Cancer Detection and Cancer Prevention'. This database was used to extract a data set with 136 attributes and 859 records. The class attribute for these experiments was the result of *H. pylori* test (positive or negative), which was carried out for each participant in the study. The selected attributes describe features of the individuals like nationality, education level, family situation, income level, occupation, smoking and alcohol consumption habits, physical activity, lifestyle, working environment, eating habits, family members, medical history, as well as age, gender, weight and height.

The experiments were carried out using *OrangeCanvas* data analysis tool with a unified experiment process model. This model provided attribute informativeness evaluation based on *CfsSubsetEval* method, using *genetic algorithm* based search approach. Classification was carried out using *C4.5*, *CN2* and *kNN* algorithms, evaluating the classifiers by *overall accuracy*, *sensitivity* and *specificity*. Classification results were evaluated using *10-fold cross-validation*.

The initial data set with 136 attributes obtained from the database was processed in order to increase its quality by

transforming, aggregating, merging and selecting the most informative results, leaving a data set with 65 attributes.

The experiments were carried out according to the process whose diagram is displayed in Fig. 3.

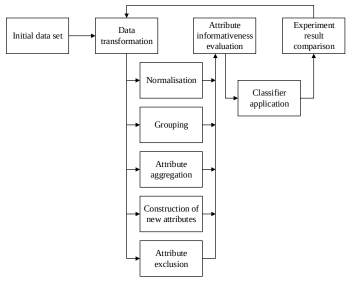


Fig. 3. Process of experiment execution.

The data set transformation involved several data preprocessing methods. Then the data set obtained in the preprocessing stage was processed by evaluating attribute informativeness and removing the least informative attributes. Then the data set with the determined most informative attributes was used in classification applying several classification algorithms. The obtained results were logged and mutually compared based on overall accuracy, sensitivity and specificity of the classifiers. These metrics were calculated from confusion matrices representing results of the various classifiers.

One of the attribute transformation processes is presented in Table I, where attributes *Height* and *Weight* were used to calculate and construct a new attribute *BMI*, which was used to aggregate a nominal attribute *BMI Categories*.

TABLE I
DATA TRANSFORMATION RESULTS

Height	Weight	BMI	<b>BMI</b> Categories			
175	96	31	Obesity			
175	108	35	Obesity			
174	66	22	Normal weight			
168	91	32	Obesity			
185	103	30	Obesity			
187	83	24	Normal weight			
190	86	24	Normal weight			
162	77	29	Overweight			

Table II shows the process of several attribute grouping to derive a new attribute characterising *consumption intensity of smoked food: days per week* and *months per year*.

Table III shows experiment results acquired using different methods of data transformations. After the 3rd experiment, the number of data set records was decreased by removing respondents who had used medicine to eradicate infection caused by *H. pylori*.

TABLE II
DATA GROUPING RESULTS

Days per week	week Months per Consumption of year smoked food			
0	0	0	None	
1–2	1-4	1	Low	
3–5	4–7	2	Medium	
6–7	8-12	3	High	

Further experiments were also carried out without the attributes that described the use of medicine for *H. pylori* eradication.

The best result of all experiments is the one achieved in the 16th experiment, which used C4.5 classification algorithm overall classifier accuracy of 68 %, sensitivity of 85 % and 29 % of specificity. This result was obtained in a data set with the following 20 attributes: Age, Gender, Height, Weight, Nationality, Heavy metals, Volatile chemical substances, Organic chemical substances, Chemicals used in agriculture, Vibration, Radiation, Biological factors, Smoking, Alcohol consumption, Walking, Moderate activity, Workload, Salty food, Smoked food and Diseases. Analysis of these results shows that development of H. pylori infection is promoted by hazardous habits described in literature previously: smoking, alcohol consumption, medical history and increased consumption of salt. Other factors determined in this research are the following: work environment and level of physical activities. The experiments show that hazardous work environment largely contributes to developing H. pylori infection. A significant part of attributes (38%) left in the final set are connected to the working environment. The final data set also holds the attribute Workload, which means that it also contributes to the risk of developing H. pylori infection, mainly due to the caused stress and lifestyle changes.

In order to make the results more demonstrative, relationships among attributes and respondents with positive H. pylori tests were calculated. Part of this information is presented in Fig. 4. It shows that 42 % of the respondents with positive tests are in the age range of 40 to 50 years, 25 % of the respondents regularly face volatile chemical compounds at work and 27 % face vibration at work. 41 % of these respondents smoke, 38 % of these respondents do not carry out any physical activities, 28 % have high workload.

Most of the respondents with positive test results (75 %) consume alcohol and 23 % of these respondents have had gastric diseases. These statistics leads to a conclusion that one of the highest impacts on developing *H. pylori* infection is from hazardous factors in the work environment, such as volatile chemical compounds and vibration. A large number of respondents use alcohol, which is a known and previously poven factor in *H. pylori* infection development. A little less than half of the respondents are smokers (41 %) and physically passive(39 %).

Data set	No. of	No. of No. of records	C4.5		kNN			CN2			
Data set	attributes		Acc.	Sen.	Spec.	Acc.	Sen.	Spec.	Acc.	Sen.	Spec.
Initial data set	65	859	0.62	0.79	0.28	0.54	0.61	0.42	0.64	0.90	0.16
1. Experiment	64	859	0.60	0.75	0.30	0.57	0.71	0.30	0.65	0.20	0.90
2. Experiment	56	859	0.60	0.75	0.30	0.55	0.69	0.28	0.63	0.87	0.24
3. Experiment	56	859	0.61	0.71	0.40	0.55	0.67	0.28	0.64	0.86	0.19
4. Experiment	50	742	0.62	0.72	0.42	0.55	0.68	0.30	0.63	0.86	0.16
5. Experiment	47	742	0.62	0.78	0.21	0.57	0.70	0.26	0.66	0.90	0.09
6. Experiment	27	742	0.61	0.79	0.17	0.57	0.67	0.31	0.65	0.88	0.10
7. Experiment	24	742	0.65	0.84	0.20	0.56	0.68	0.27	0.66	0.91	0.05
8. Experiment	9	742	0.70	0.96	0.07	0.65	0.78	0.32	0.71	0.98	0.04
9. Experiment	43	742	0.64	0.80	0.32	0.60	0.73	0.33	0.67	0.95	0.11
10. Experiment	40	742	0.64	0.82	0.22	0.63	0.77	0.27	0.70	0.95	0.07
11. Experiment	35	742	0.68	0.88	0.21	0.59	0.74	0.32	0.63	0.83	0.13
12. Experiment	30	742	0.66	0.81	0.27	0.62	0.77	0.25	0.65	0.87	0.12
13. Experiment	28	742	0.65	0.81	0.28	0.62	0.62	0.22	0.64	0.86	0.13
14. Experiment	26	742	0.67	0.85	0.25	0.65	0.78	0.33	0.64	0.87	0.06
15. Experiment	23	742	0.67	0.82	0.31	0.60	0.75	0.24	0.66	0.90	0.10
16. Experiment	20	742	0.68	0.85	0.29	0.63	0.77	0.27	0.65	0.90	0.04

TABLE III Experiment Results

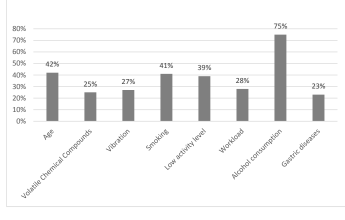


Fig. 4. Analysis of the obtained results.

## V. CONCLUSION

Experimental evaluation of the initial data set led to decreasing of the data set to 20 attributes and 742 records. Most of the experiments were carried out using data preprocessing methods and constructing different variations of the data set to determine the most suitable attributes or attribute subsets.

The experimental analysis of factor sets and data set sizes showed that these changes had mostly small impact on overall accuracy of the classifiers. The overall accuracy for C4.5varied in the range from 60 % to 70 %. Most of the variations were due to strong variations in sensitivity and specificity. Since the goal of the research was to determine factors that influence positive class, sensitivity was given the major importance but the desired specificity should be around 30 % or one correct prediction among three to avoid assigning all individuals to the positive class and receiving 0 % specificity and no useful information from the model.

The experimental evaluation determined the most suitable classifier, which was constructed using C4.5 classification algorithm. This classifier showed that an infection risk was influenced by factors contributing to *H. pylori* development like *working environment* and *physical activity*. Closer analysis of these factors showed that respondents, who had positive *H. pylori* test results, worked in an environment where they faced either *vibration* or *volatile chemical compounds*, as well as heavy *workload*. Another contributing factor was *lack of physical activity*, which was confirmed in 38 % positive respondents who did not carry out any physical exercise.

In cases when a person faces hazardous work, environment and volatile chemical compounds at work, a doctor should appoint them to *H. pylori* test, which could facilitate early detection and prevention of gastric diseases and cancer.

#### ACKNOWLEDGMENT

The present research has been developed by the GISTAR within the project "Interdisciplinary Research Group for Early Cancer Detection and Cancer Prevention" and supported by the International Agency for Research on Cancer (IARC).

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