Assessment of the Risk Factors of Coronary Heart Events Based on Data Mining With Decision Trees

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***Abstract*—Coronary heart disease (CHD) is one of the major causes of disability in adults as well as one of the main causes of death in the developed countries. Although significant progress has been made in the diagnosis and treatment of CHD, further inves- tigation is still needed. The objective of this study was to develop a data-mining system for the assessment of heart event-related risk factors targeting in the reduction of CHD events. The risk factors investigated were: 1) before the event: a) nonmodifiable—age, sex, and family history for premature CHD, b) modifiable—smoking before the event, history of hypertension, and history of diabetes; and 2) after the event: modifiable—smoking after the event, systolic blood pressure, diastolic blood pressure, total cholesterol, high- density lipoprotein, low-density lipoprotein, triglycerides, and glu- cose. The events investigated were: myocardial infarction (MI), percutaneous coronary intervention (PCI), and coronary artery bypass graft surgery (CABG). A total of 528 cases were collected from the Paphos district in Cyprus, most of them with more than one event. Data-mining analysis was carried out using the C4.5 decision tree algorithm for the aforementioned three events using five different splitting criteria. The most important risk factors, as extracted from the classification rules analysis were: 1) for MI, age, smoking, and history of hypertension; 2) for PCI, family his- tory, history of hypertension, and history of diabetes; and 3) for CABG, age, history of hypertension, and smoking. Most of these risk factors were also extracted by other investigators. The highest percentages of correct classifications achieved were 66%, 75%, and 75% for the MI, PCI, and CABG models, respectively. It is antici- pated that data mining could help in the identification of high and low risk subgroups of subjects, a decisive factor for the selection of therapy, i.e., medical or surgical. However, further investigation with larger datasets is still needed.**

***Index Terms*—Coronary heart disease (CHD), data mining, de- cision trees, risk factors.**

1. INTRODUCTION

**C**

ORONARY heart disease (CHD) is the single most com- mon cause of death in Europe, responsible for nearly two million deaths a year [1]. Advances in the field of medicine over the past few decades enabled the identification of risk factors that may contribute toward the development of CHD. However,

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this knowledge has not yet helped in the significant reduction of CHD incidence. There are several factors that contribute to the development of a coronary heart event. These risk fac- tors may be classified into two categories, not modifiable and modifiable. The first category includes factors that cannot be altered by intervention such as age, gender, operations, family history, and genetic attributes [2]–[4]. Modifiable risk factors are those for which either treatment is available or in which alternations in behavior can reduce the proportion of the pop- ulation exposed. Established, modifiable risk factors for CHD currently include smoking, hypertension, diabetes, cholesterol, high-density lipoprotein, low-density lipoprotein, triglycerides [5], [6].

The objective of this study was to develop a data mining system based on decision trees for the assessment of CHD- related risk factors targeting in the reduction of CHD events. Data-mining analysis was carried out using the C4.5 decision tree algorithm using five different splitting criteria for extract- ing rules based on the aforementioned risk factors. Preliminary results of this study were previously published [7].

Many studies have been carried out investigating CHD and re- lated risk factors. Some of them used the Framingham equation to describe the population in a region or country [8], [9], whereas other studies examined the features of available Framingham- based risk calculation [10]. The American Heart Association (AHA) assessed multiple risk factors and also developed new guidelines for CHD [11], [12]. Furthermore, results from the European Action on Secondary and Primary Prevention by In- tervention to Reduce Events (EUROASPIRE) revealed the im- portant risk factors through various surveys across European countries [2]–[4].

Data mining facilitates data exploration using data analysis methods with sophisticated algorithms in order to discover un- known patterns. Such algorithms include decision trees that have been used extensively in medicine. According to Podgorelec *et al.* [13] decision-tree-based algorithms give reliable and ef- fective results that provide high-classification accuracy with a simple representation of gathered knowledge, and are es- pecially appropriate to support decision-making processes in medicine.

Several studies have been carried out that investigated the usefulness of decision tree models in CHD-related problems. Ordonez [14], [15] investigated decision trees and association rules to predict CHD based on the risk factors sex, smok- ing, cholesterol, and age. Gamberger *et al.* [16] used a deci- sion support method to target high-risk groups for CHD using risk factors like smoking, cholesterol and hypertension. Tsien

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*et al.* [17] used also classification trees and logistic regression building three different models for myocardial infarction (MI) and examining also the significance of these models. Decision- tree-based software tools were developed in [18] and [19] to aid in the diagnosis of CHD. Rao *et al.* [18] presented a framework to create structured clinical data for CHD. Zavrsnik *et al.* [19] used decision trees and created the ROSE tool for use in car- diology. Furthermore, Polat *et al.* [20] developed decision tree based models for the classification of CHD, achieving a correct classification score of 82%. Moreover, Pavlopoulos *et al.* [21] used the C4.5 algorithm decision trees to analyze the different heart sound features, which assist clinicians to make a better diagnosis in CHD.

Several other studies investigated different technologies for the assessment of CHD, including logistic regression [17], as- sociation rules [7], [15], fuzzy modeling [22], [23], neural net- works [24], and other.

In this study, we investigate how data mining based on deci- sion trees can help for the evaluation of the risk of CHD. The aim is to identify the most important risk factors based on the classification rules to be extracted. These rules will enable the better management of the patient targeting in the reduction of events, as well as, reduction of the cost of therapy, due to the expected restriction of interventions in necessary cases only.

The rest of the paper is organized as follows. Section II de- scribes the material and methods, section III the results and Section IV the discussion.

1. MATERIALS AND METHODS
2. *Data Collection, Cleaning, and Coding*

Data from 1500 consecutive CHD subjects were collected be- tween the years 2003–2006 and 2009 (300 subjects each year) according to a prespecified protocol, under the supervision of the participating cardiologist (Dr. J. Moutiris, second author of this paper) at the Department of Cardiology, at the Paphos General Hospital in Cyprus. Subjects had at least one of the following criteria on enrollment, history of MI, or percutaneous coronary intervention (PCI), or coronary artery bypass graft surgery (CABG). Data for each subject were collected as given in Table I: 1) risk factors before the event, a) nonmodifiable— age, sex, and family history (FH); 2) modifiable—smoking be- fore the event (SMBEF), history of hypertension (HxHTN), and history of diabetes (HxDM); and 2) risk factors after the event, modifiable—smoking after the event (SMAFT), sys- tolic blood pressure (SBP) in mmHg, diastolic blood pressure (DBP) in mmHg, total cholesterol (TC) in mg/dL, high-density lipoprotein (HDL) in mg/dL, low-density lipoprotein (LDL) in mg/dL, triglycerides (TG) in mg/dL, and glucose (GLU) in mg/dL.

To clean the data, the fields were identified, duplications were extracted, missing values were filled, and the data were coded as given in Table I. After data cleaning, the number of cases was reduced as given in Table II, mainly due to unavailability of biochemical results.

TABLE I CODING OF RISK FACTORS

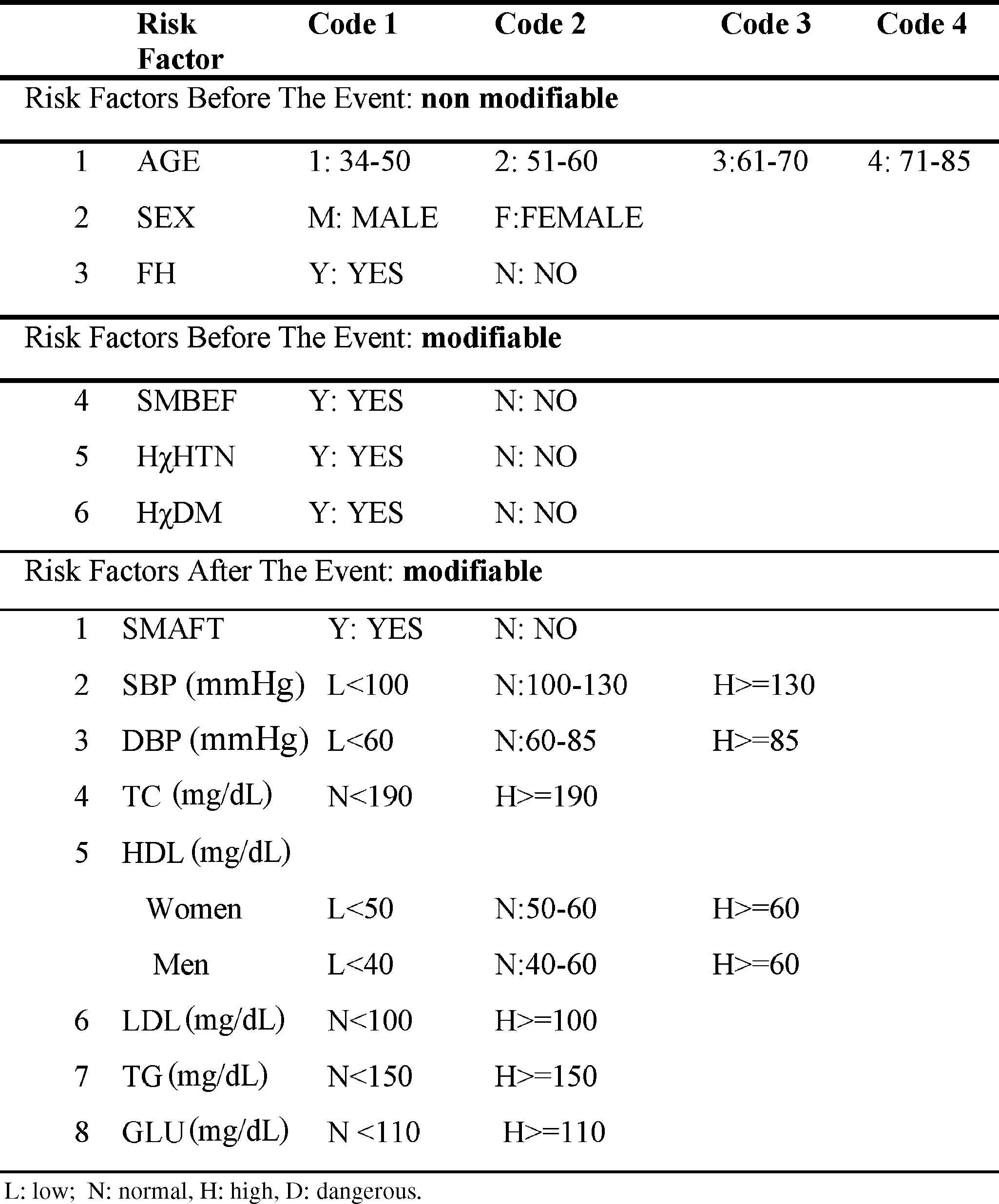
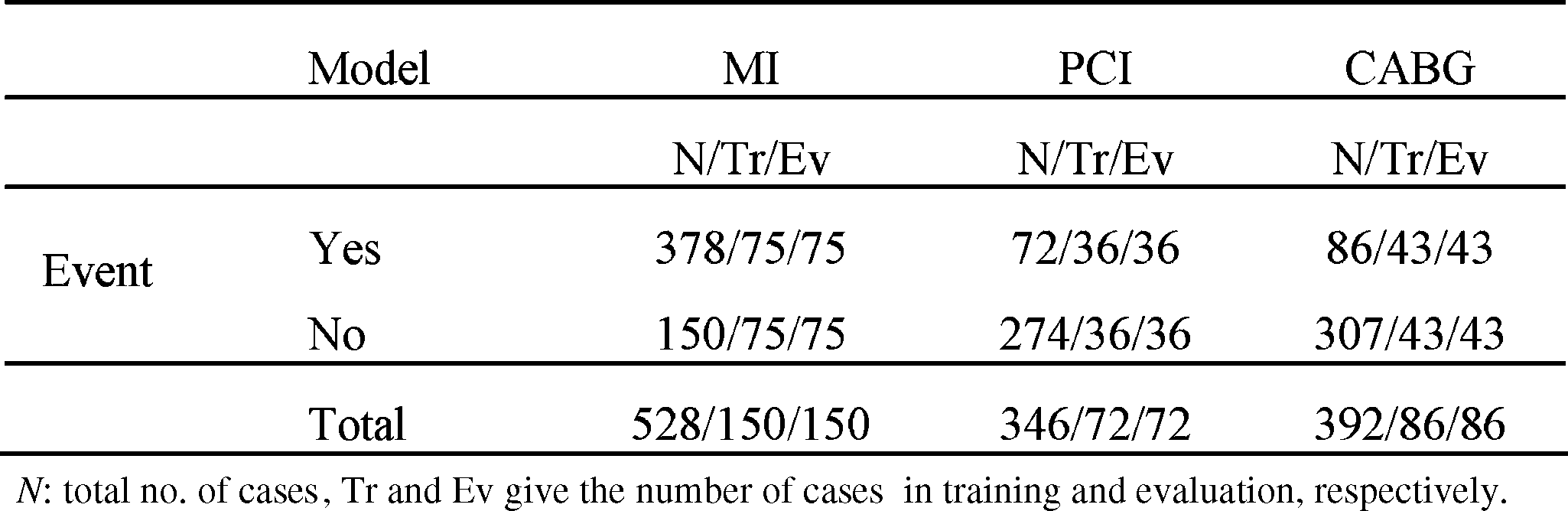


TABLE II

NO. OF CASES PER SET OF RULES/MODELS INVESTIGATED



1. *Classification by Decision Trees*

The C4.5 algorithm [25], which uses the divide-and-conquer approach to decision tree induction, was employed. The al- gorithm uses a selected criterion to build the tree. It works top–down, seeking at each stage an attribute to split on that which best separates the classes, and then recursively process- ing the sub problems that result from the split. The algorithm uses heuristics for pruning derived based on the statistical sig- nificance of splits.

*Algorithm Generate Decision Tree [25], [26]: Input:*

* 1. Training dataset *D*, which is a set of training observations and their associated class value.
  2. Attribute list *A*, the set of candidate attributes.
  3. Selected splitting criteria method.

*Output:* A decision tree.

*Method:*

1. Create a node *Nd*.
2. If all observations in the training dataset have the same class output value *C*, then return *Nd* as a leaf node labeled with *C*.
3. If attribute list is empty, then return *Nd* as leaf node labeled with majority class output value in training dataset.
4. Apply selected splitting criteria method to training dataset in order to find the ‘‘best” splitting criterion attribute.
5. Label node *Nd* with the splitting criterion attribute.
6. Remove the splitting criterion attribute from the attribute list.
7. For each value *j* in the splitting criterion attribute.
   1. Let *Dj* be the observations in training dataset satis- fying attribute value *j*.
   2. If *Dj* is empty (no observations), then attach a leaf node labeled with the majority class output value to node *Nd*.
   3. Else attach the node returned by generate deci-

sion tree (*Dj* , attribute list, selected splitting criteria method) to node *Nd*.

1. *Gini Index (GI):* The Gini index is an impurity-based cri- terion that measures the divergence between the probability dis- tributions of the target attributes values [28]

*v*

Σ*− ×*

GiniIndex(*D*) = Gini(*D*) *pj* Gini(*Dj* ) (2.4)

*j* =1

*m*

ΣGini (*D*) = 1 *− p .* (2.5)

2

*i*

*i*=1

1. *Likelihood Ratio Chi-Squared Statistics (χ*2 *):* The like- lihood ratio chi-squared statistic is useful for measuring the statistical significance of the information gain criterion [29]

*G*2 (*A, D*) = 2 *×* ln (2) *× |D|×* InfoGain (*A*) *.* (2.6)

1. *Gain Ratio (GR):* Gain ratio biases the decision tree against considering attributes with a large number of distinct values. So it solves the drawback of information gain [25]
2. End for.
3. Return node *Nd*.

In this study, the following splitting criteria were investigated

that are briefly presented shortly: information gain, gini index,

GainRatio (*A*) = InfoGain(*A*)

SplitInfo*A* (*D*)

*v*

Σ*− ×*

(2.7)

likelihood ratio chi-squared statistics, gain ratio, and distance measure.

SplitInfo*A*

1. = *|Dj |* log2

*j* =1 *|D|*

*|Dj | .* (2.8)

*|D|*

* 1. *Information Gain (IG):* Information gain is based on Claude Shannon’s work on information theory. InfoGain of an attribute *A* is used to select the best splitting criterion attribute. The highest InfoGain is selected to build the decision tree [27]

*5) Distance Measure (DM):* Distance measure, like GR, nor- malizes the impurity criterion (GI). It suggests normalizing it in a different way [30]

InfoGain(*A*) = Info (*D*) *−* Info

(*D*) (2.1)

DM (

) = Gini (*D*) *.* (2.9)

*A A v*

Σ*−*

Σ

*j* =1

*m i*=1

*pij*

*×* log2

(*pij* )

where *A* is the attribute investigated.

*m*

Σ*−*

Info (*D*) = *pi* log2 (*pi*) (2.2)

*i*=1

where

*pi* = probability(class *i* in dataset *D*);

A data-mining tool was developed by our group that supports the C4.5 decision tree algorithm using the aforementioned cri- teria. Overfitting is a significant practical difficulty for decision tree learning. Therefore, pruning is implemented to avoid over- fitting. We implemented the bottom-up pruning algorithm using Laplace error estimation. While the decision tree is built and a leaf node is created, then the Laplace error [31] is estimated as follows:

*m* = number of class values*.*

*v*

where

*E* (*D*) = *N − n* + *m −* 1

*N* + *m*

(2.10)

Info*A* (*D*) = Σ *| |* Info (*D* ) (2.3)

*D j*

*j*

where

*j* =1

*|D|*

*C* = class value majority class in *D*;

*N* = number of observations in *D*;

*Dj* = number of observations with attribute value *j* in dataset *D*;

*| |*

*D* = total number of observations in dataset *D*;

*| |*

*Dj* = sub dataset of *D* that contains attribute value *j*; *v* = all attribute values.

Although information gain is usually a good measure for deciding the relevance of an attribute, it is not perfect. A problem occurs when information gain is applied to attributes that can take on a large number of distinct values. When that happens, then gain ratio is used instead.

*n* = number of observations has class value *C.*

As the algorithm returns to the root node, the error of the leaf node is passed to the father node. The father node calculates the total error of all of its children and its own error. If the father’s error is less than the total error of the children, then the father node is pruned and replaced by a leaf node with the majority class value. If the father’s error is greater than the total error of the children, then no more pruning is done to the path and the returned error is zero.

1. *Classification Models Investigated*

The following sets of models were investigated as given in Table II.

* 1. MI: MI versus non-MI. Subjects having myocardial infrac- tion were marked as symptomatic and the rest as asymp- tomatic.
  2. PCI: PCI versus non-PCI. Subjects having only PCI were marked as symptomatic and the rest as asymptomatic. Subjects having both PCI and MI were excluded.
  3. CABG: CABG versus non-CABG. Subjects having only CABG were marked as symptomatic and the rest as asymptomatic. Subjects having both CABG and MI were excluded.

For each set of models, three different subsets of runs were carried out as given in the following:

1. with risk factors before the event (B);
2. with risk factors after the event (A); and
3. with risk factors before and after the event (B + A).

For each model, for each splitting criterion, 20 runs were carried out with random sampling [32] of equal number of cases used for training and evaluation as given in Table II. A total of 300 runs were carried out for each set of models [i.e., 20 runs 5 splitting criteria 3 (for B, A, and B +A datasets)].

*× ×*

The Wilcoxon rank sum test [33] was also carried out to investigate if there was or not significant difference between the five splitting criteria used as well as between the B, A, and B + A decision tree models at *p <* 0.05.

1. *Performance Measures*

In order to evaluate the performance of our results we used the following measures [34].

* 1. *Correct classifications* (%CC): is the percentage of the correctly classified records; equals to (TP + TN)/*N*.
  2. *True positive rate* (%TP): corresponds to the number of positive examples correctly predicted by the classification model.
  3. *False positive rate* (%FP): corresponds to the number of negative examples wrongly predicted as positive by the classification model.
  4. *True negative rate* (%TN): corresponds to the number of negative examples correctly predicted by the classification model.
  5. *False negative rate* (%FN): corresponds to the number of positive examples wrongly predicted as negative by the classification model.
  6. *Sensitivity*: is defined as the fraction of positive examples predicted correctly by the model, equals to TP/(TP + FN).
  7. *Specificity*: is defined as the fraction of negative examples predicted correctly by the model, equals to TN/(TN + FP).
  8. *Support*: is the number of cases for which the rule applies (or predicts correctly; i.e., if we have the rule *X Z*, Support is the probability that a transaction contains

*→*

*{X*, *Z}* [26]

Support = *P* (*XZ*) = no of cases that satify *X* and *Z .*

*|D|*

* 1. *Confidence*: is the number of cases for which the rule applies (or predicts correctly), expressed as a percentage of all instances to which it applies (i.e., if we have the rule *X Z*, Confidence is the conditional probability that a transaction having *X* also contains *Z*) [26]

Conﬁdence = *P* (*Z|X*) = *P* (*XZ*) *.*

*→*

*P* (*X*)

1. *Calculation of the Risk*

For each subject, we used the Framingham equation [8]–[10] to calculate the risk for an event to occur. We separated the subjects into two categories, those who have had an event and those who have not had an event. Then, for each extracted rule, we found out the subjects matching that rule and computed the average event risk per rule based on the risk value of each subject (see last two columns of Table V). It is noted that values of risk lower than 5%, between 5–10%, and higher than 10% classify a subject as low, intermediate, and high risk, respectively.

1. RESULTS

Table III tabulates the classification results of the three set of models investigated for the five different splitting criteria using risk factors before the event (B), after the event (A), and before and after (B + A). The median (Me), minimum (*m*), and maximum (*M*) for 20 runs are given for %CC, %TP, and

%FP, whereas for sensitivity and specificity only the median values are given. Table IV gives the three most important risk factors obtained from the classification decision tree models. Also, selected rules of the models obtained in Table III are given in Table V as well as the risk per rule computed using the Framingham equation.

1. *MI Models*

There was no significant difference for the different splitting criteria investigated for %CC using the Wilcoxon rank sum test at *p <* 0.05. As shown in Table III, comparable performance in the region of 60% for %CC was obtained for the B, A, and B + A risk factor models for all splitting criteria. Better perfor- mance was obtained for the B + A models, where the median of the %CC ranged from 62% to 63%, respectively. The best model was obtained when using the GI splitting criterion for the B + A risk factor codings with a maximum %CC = 66%.

The most important risk factors as given in Table IV were for the B models, age, history of hypertension, and smoking before the event, for the A models, systolic blood pressure, smoking after the event, and diastolic blood pressure, and for the B + A models, age, systolic blood pressure, smoking, and history of hypertension.

Based on the decision tree model, sample rules could be extracted. For example, as given in Table IV:

*Rule 1.3 and 1.4:*

* 1. The percentage of subjects aged 51–60 with history of hypertension who are non smokers and have event is

TABLE III

CLASSIFICATION RESULTS OF THE THREE SET OF MODELS INVESTIGATED FOR THE FIVE DIFFERENT SPLITTING CRITERIA USING RISK FACTORS BEFORE THE

EVENT (B), AFTER THE EVENT (A), AND BEFORE AND AFTER (B+A)

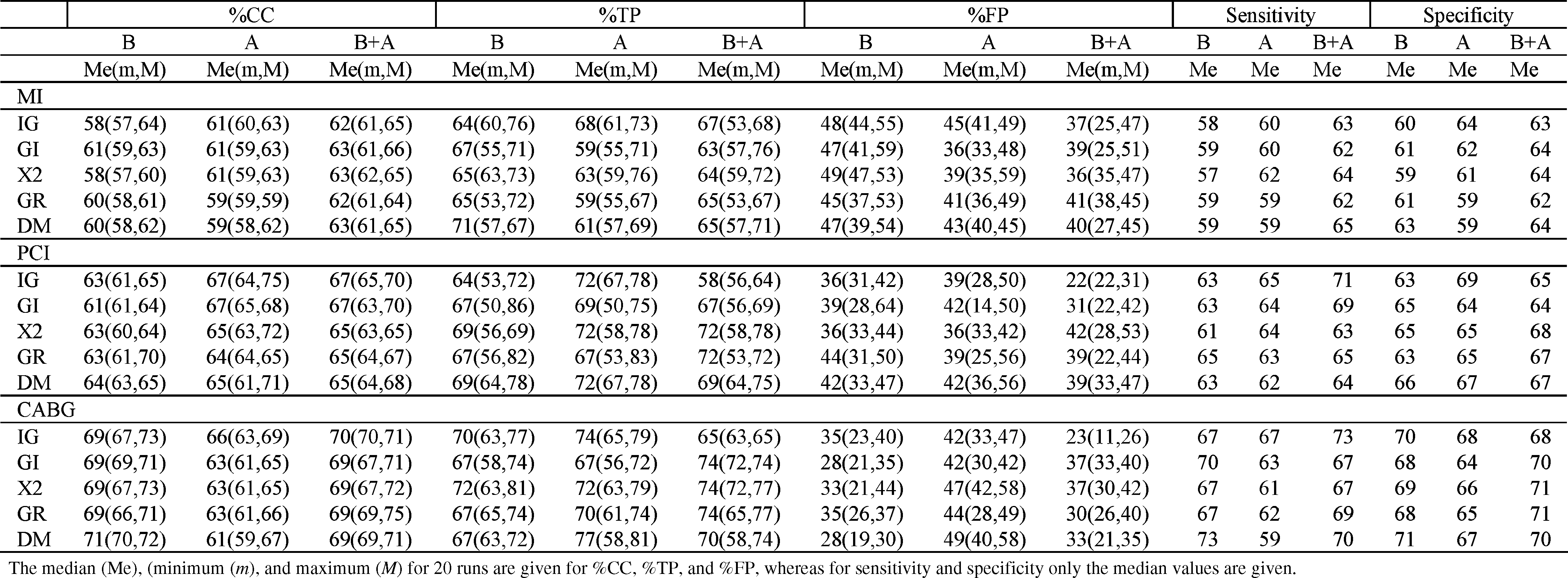
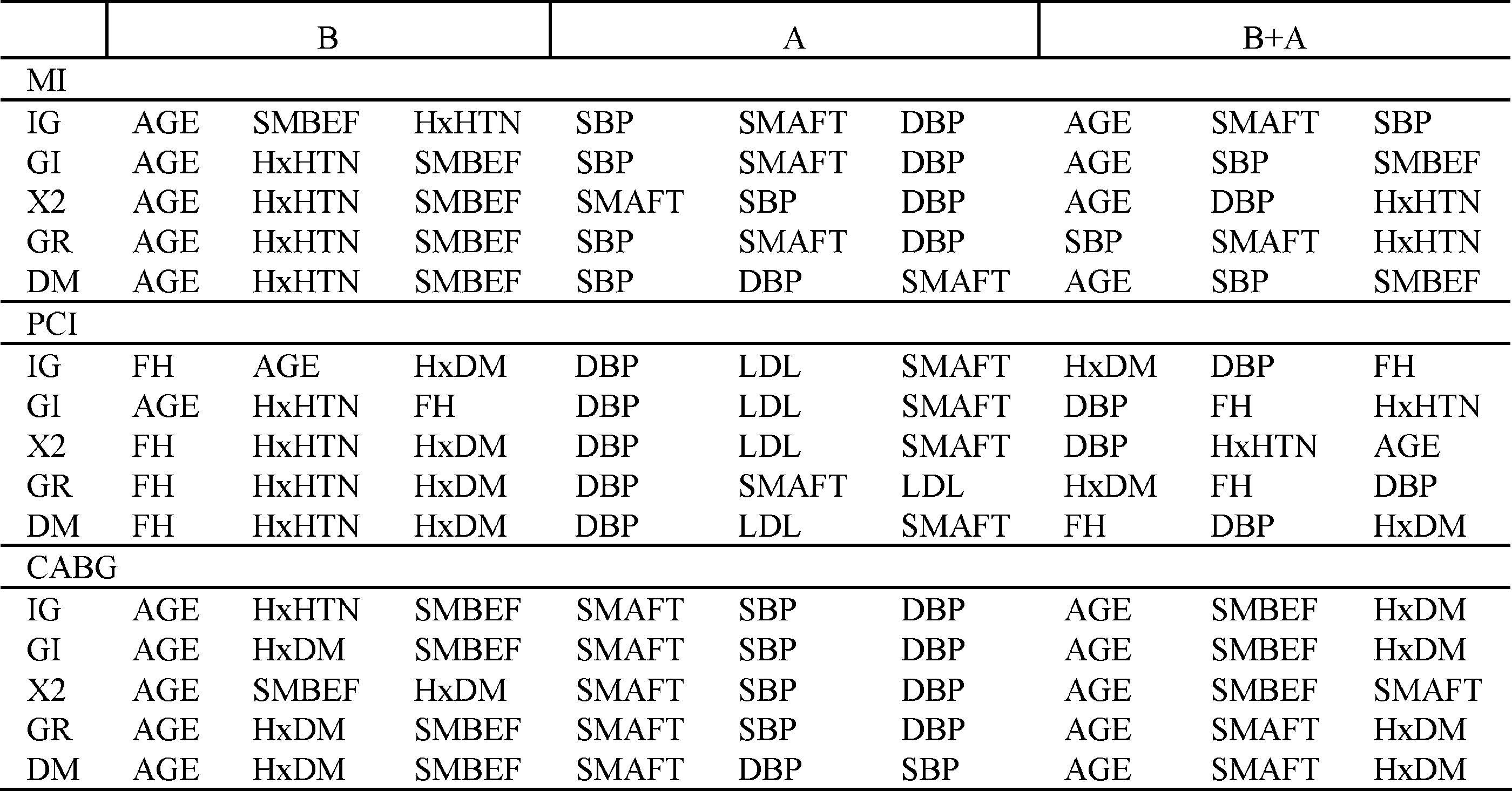


TABLE IV

THREE MOST IMPORTANT RISK FACTORS OF THE THREE SET OF MODELS INVESTIGATED GIVEN IN TABLE III FOR THE FIVE DIFFERENT SPLITTING CRITERIA

USING RISK FACTORS BEFORE THE EVENT (B), AFTER THE EVENT (A), AND BEFORE AND AFTER (B + A)



approximately the same with those who were smokers and did not have an episode.

For the MI models, there were 0/0 (0/0%), 28/7 (5.3/1.3%), and 330/163 (62.5/30.9%) subjects with event yes/no, with low, intermediate, and high risk, respectively. Moreover, the average event risk per rule ranged from 11.8% to 15.0%, i.e., all rules were classified as high risk (see Table IV). Also, there was no difference between the rule event risk for an MI event to occur versus not to occur.

1. *PCI Models*

For the PCI models, slightly better performance was obtained compared to the MI models. Better performance was obtained for the A and B + A models, with the median of %CC ranging from 65% to 67%. Again, similar performance was obtained for all splitting criteria with no significant difference.

The most important risk factors were for the before risk fac- tors models, age, family history, history of hypertension and history of diabetes, for the after risk factors models, diastolic blood pressure, low density lipoprotein, and smoking after the event, and for the before and after risk factors models, history of diabetes, diastolic blood pressure, family history, history of hypertension, and age.

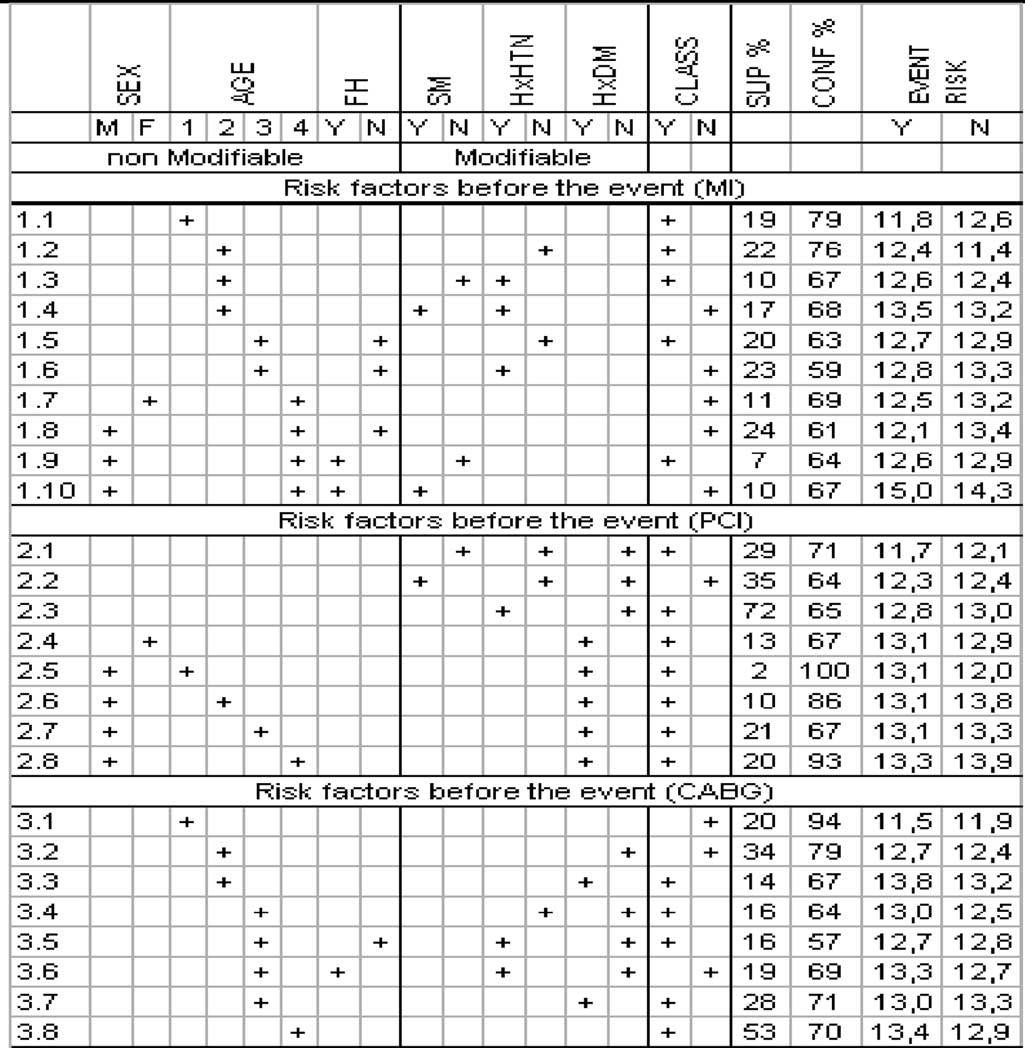
Based on the rules given in Table IV:

Rules 2.5–2.8 for diabetes subjects the number of PCI events increase with age (support increases from 2% to 20%).

For the PCI models, there were 0/0 (0/0%), 20/15 (3.8/2.8%), and 193/300 (36.6/56.8%) subjects with event yes/no, with low, intermediate, and high risk respectively. The average event risk per rule ranged from 11.7 to 13.9%, i.e all rules were classi- fied as high risk (see Table IV). Also, there was no difference between the rule event risk for a PCI event to occur vs not to occur.

TABLE V

SELECTED RULES FROM MODELS GIVEN IN TABLE III (BASED ON THE CODING OF THE RISK FACTORS GIVEN IN TABLE II)



1. *CABG Models*

Highest performance was obtained for the CABG models, with median of %CC in the region of 70%. As in the aforemen- tioned two set of models, there was no significant difference in the models obtained with the different splitting criteria. The highest performance was obtained for the GR splitting criterion, for the B + A model, where the maximum %CC = 75%.

The most important risk factors based on Table IV were for the before risk factors models, age, history of hypertension, history of diabetes, and smoking before the event, for the after risk factors models, smoking after the event, systolic blood pressure, and diastolic blood pressure, and for the before and after risk factors models, age, smoking before the event, smoking after the event, and history of diabetes. Based on the rules given in Table V:

*Rules 3.2 and 3.3*:

* 1. CABG occurs usually in subjects aged between 51 and 60 years old when they have history of diabetes.

*Rules 3.5 and 3.6:*

1) Family history is not an important risk factor for CABG. For the CABG models, there were 0/0 (0/0%), 9/26 (1.7/4.9%), and 206/287 (39/54.4%) subjects with event yes/no, with low, intermediate, and high risk respectively. Similar to the previous two models, the average event risk per rule varied very little, ranging from 11.5 to 13.3%, and all rules were classified as high risk (see Table IV). Also, there was no difference be- tween the rule event risk for a CABG event to occur vs not to

occur.

1. DISCUSSION

The events investigated through this study were: MI, PCI, and CABG. Three classification models were developed based on decision trees for classifying MI, PCI, and CABG patients, where the highest percentage of correct classifications obtained were 66%, 75%, and 75%, respectively. Although different risk factors were obtained for the MI, PCI, and CABG models in- vestigated, the most important risk factors, as extracted from the classification rule analysis were: sex, age, smoking, blood pressure, and cholesterol. It is important to note that the latter three risk factors can be modified; therefore the CHD risk of a subject may be reduced through a proper control of these fac- tors. Furthermore, the importance of smoking in increased CHD risk was clearly illustrated.

The above findings and risk factors were also extracted by other investigators [35]. The EUROASPIRE study with EUROASPIRE surveys (I, II, III) involved various European populations and also included additional risk factors such as obesity. All Euroaspire surveys were reviewed together and combined results were extracted [4]. A general outcome was the fact that patients do not follow the advice and recommenda- tions of their physicians. In comparison with the EUROASPIRE survey, our findings concerning the modifiable risk factors after the event are the following [4]:

1. 14% of subjects smoke after the event (16% in EUROASPIRE);
2. 22% of subjects had high blood pressure (26% in EUROASPIRE);
3. 34% of subjects had high total cholesterol (31% in EUROASPIRE); and
4. 45% of subjects had low-density lipoprotein (31% in EUROASPIRE).

In the EUROASPIRE survey, smoking, blood pressure, and cholesterol were found to be important risk factors [2], [4]. It was concluded that wide variations exist between 15 countries in the risk factor prevalence’s and the use of cardioprotective drug therapies [3]. Also, there is still considerable potential throughout Europe to raise standards of preventive care in order to reduce the risk of recurrent disease and death in patients with CHD.

Furthermore, additional observations that could be extracted from the database investigated in this study regarding the non- modifiable risk factors in comparison with EUROASPIRE sur- vey [4] are the following:

1. 14% of subjects were female (24.7% in EUROASPIRE);
2. 9% of subjects were 50 years old (23.1% in EUROASPIRE);

*≤*

1. 28% were between 51 and 60 years old (33.8% in EUROASPIRE);
2. 39% of subjects were between 61 and 70 years old (43.1% in EUROASPIRE); and
3. 24% of subjects were between 71 and 84 years old.

No female subject was under the age of 50 years old; only male subjects were found under this age.

Rea *et al.* [35] concluded that smoking was associated with an elevated risk for recurrent coronary events, whereas Gamberger

*et al.* [16] mention the relationship between the risk factors cholesterol and overweight.

Wang *et al.* [8] used the risk factors age, sex, cholesterol, HDL, blood pressure, diabetes, and smoking to predict CHD. They used the Framingham function and concluded that the traditional risk factors have different degrees of impact and/or than other factors are contributing to risk.

It should be noted that the results of our study based on a small city in the island of Cyprus are comparable with other studies, as it is known that traditional risk factors have different degrees of impact and/or that other factors are contributing to risk. A population-specific risk function is needed as also indicated by other investigators [8].

The values of risk computed for each subject were between 7% and 15.5% that fall into the range of none (0%) for low risk, 35 (6.6%) for intermediate risk, and 493 (93.4%) for high risk. Although an average rule risk was computed for each rule, the values extracted for an event to occur or not are very close, not making possible the differentiation between high and low risk subgroups of subjects. This finding should somehow be expected, given that almost all of the subjects used for deriving the proposed models fall into the high, risk group. Thus, the proposed methodology should be further investigated by using a more heterogenous group of subjects, covering numerous cases

of low and medium risk.

Ordonez [14] using the C4.5 decision tree algorithm and as- sociation rules for the prediction of cardiac disease based on 25 risk factors documented that association rules generally include simpler predictive rules than decision tree rules [15]. The use- fulness of association rules in the analysis of CHD risk factors was also investigated by our group on a similar database with this study [36]. The results regarding the most important risk factors were similar.

Tsien *et al.* [17] in their study indicated that classification trees, which have certain advantages over logistic regression models, may perform similar to logistic regression models in the diagnosis of patients with MI.

The following five different criteria were investigated, in- formation gain, gini index, likelihood ratio chi-squared statis- tics, gain ratio, and distance measure, that resulted in models with similar performance, with no significant difference be- tween them. Thus any one of the splitting criteria investigated could be used for the datasets in this study. This finding is in agreement with this study for developing the decision tree mod- els, that documented that the choice of splitting criteria does not make much difference on the tree performance [24], [32]. Also, the different splitting criteria, agreed on the most im- portant risk factors. To the best of our knowledge no simi- lar study was found in the literature comparing the five dif- ferent criteria investigated in this study for the problem of CHD.

Concluding, comparing our findings with other studies:

1) a data mining system was proposed to extract rules for CHD events, 2) the rules extracted facilitated the grouping of risk factors into high and low risk factors, and 3) the rules extracted are associated with an event risk, however, this needs further investigation.

It is anticipated that data mining based on decision trees could help in the identification of risk subgroups of subjects for de- veloping future events and it might be a decisive factor for the selection of therapy, i.e., angioplasty or surgery. Moreover, the extracted models and rules could help to reduce CHD morbid- ity and possibly, mortality. However, further investigation with larger datasets and other rule extraction algorithms and criteria are still needed.

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