Developing the breast cancer risk prediction system using hybrid machine learning algorithms

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Abstract:

BACKGROUND: Breast cancer (BC) is the most common cause of cancer-related deaths in women globally. Currently, many machine learning (ML)-based predictive models have been established to assist clinicians in decision making for the prediction of BC. However, preventing risk factor formation even with having healthy lifestyle behaviors or preventing disease at early stages can significantly lead to optimal population-wide BC health. Thus, we aimed to develop a prediction model by using a genetic algorithm (GA) incorporating several ML algorithms for the prediction and early warning of BC.

MATERIAL AND METHODS: The data of 3168 healthy individuals and 1742 patient case records in the BC Registry Database in Ayatollah Taleghani hospital, Abadan, Iran were analyzed. First, a modified hybrid GA was used to perform feature selection and optimization of selected features. Then, with the use of selected features, several ML algorithms were trained to predict BC. Afterward, the performance of each model was measured in terms of accuracy, precision, sensitivity, specificity, and receiver operating characteristic (ROC) curve metrics. Finally, a clinical decision support system based on the best model was developed.

RESULTS: After performing feature selection, age, consumption of dairy products, BC family history, breast biopsy, chest X-ray, hormone therapy, alcohol consumption, being overweight, having children, and education statuses were selected as the most important features for prediction of BC. The experimental results showed that the decision tree yielded a superior performance than other ML models, with values of 99.3%, 99.5%, 98.26% for accuracy, specificity, and sensitivity, respectively.

CONCLUSION: The developed predictive system can accurately identify persons who are at elevated risk for BC and can be used as an essential clinical screening tool for the early prevention of BC and serve as an important tool for developing preventive health strategies.

Keywords: Breast cancer, lifestyle, machine learning, prevention

Introduction

Breast cancer (BC) is one of the most frequent cancers in women and the second leading cause of death after lung cancer, according to the World Health Organization (WHO) report.[1-3] BC is the most frequent malignancy among females, with an estimated 11.7% of all cancer cases and 2.3 million new cases in 2020.[4] Several biomarkers have been identified for detecting and predicting this disease.[5,6] Family history is one of the known risk factors for BC. Age is another factor that may be associated with the risk of BC. For example, studies state that women with a family history of BC and those who are more than 40 years old are highly susceptible to have the risk of BC.[7] Evidence indicates the relationship between diet habits, especially fat consumption in postmenopausal women,

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and BC. Other evidence demonstrated that hormonal factors, having a history of benign breast tumors, family history of BC, and genetic factors increase the risk of BC incidence. Obesity, which is prevalent in about 20% of the population in developed countries, is another factor that increases the risk of BC during the postmenopausal period. Diabetes is another risk factor; 5%-16% of patients with BC who are more than 65 years old have diabetes. The incidence of BC and type 2 diabetes mellitus is prevalent among older people who are a common risk factor in obese individuals. Regarding the lifestyle of the people in the modern era, weight gain and obesity are prevalent and may thus increase the risk of BC. However, the reports vary; some researchers suggest that people with a BMI of more than 30 are highly susceptible to the incidence of BC in premenopausal periods. Although various factors influence the risk of BC, these factors are not linearly associated, and the relationship between them is complicated. Therefore, using machine learning (ML) may be influential as these techniques do not consider the association of the variables nonlinear and are compatible with complicated relations. In recent years, many ML techniques have been employed for predicting and classifying BC outcomes. ML algorithms consist of supervised and unsupervised methods. We considered supervised methods. In the supervised approach, we used a part of our data as a training dataset to train our model, and then we tested the model with the part of data that is new to the algorithm.

To date, many studies have tried to predict BC by using outstanding ML techniques. For example, Dhahri et al. introduced an automated BC detection based on ML algorithms. Their study was based on GA and ML techniques; they aimed to develop a system to accurately differentiate between benign and malignant breast tumors. They applied several techniques to choose the best features and perfect parameter values as ML classifiers inputs. They found that the GA can automatically detect the best model by combining feature preprocessing methods and classifier algorithms.

Salod et al., in their study, compared the performance of ML algorithms in BC screening and detection. They used anthropometric blood analysis data from female BC patients and volunteer healthy controls of the UCI ML repository datasets. They applied eight ML algorithms, including logistic regression, support vector machine (SVM), k-nearest neighbors (KNN), decision tree (DT), random forest (RF), adaptive boosting (AdaBoost), gradient boosting machine (GBM), and eXtreme gradient boosting (XGBoost) and selected the best model considering the performance metrics of accuracy, precision, recall or sensitivity, specificity, F1 score, receiver operating characteristic (ROC) curve, and area under the ROC curve (AUC).

In another study conducted by Soltani Sarvestani et al., efficient networks for BC data mining from clinically collected datasets were investigated. By using various data mining techniques, they aimed to find out the percentage of disease development. The performance of the statistical neural network models, self-organizing map (SOM), radial basis function network (RBF), general regression neural network (GRNN), and probabilistic neural network (PNN) were evaluated on the Wisconsin Breast Cancer data (WBCD) and the Shiraz Namazi Hospital Breast Cancer Data (NHBCD). The results were considered, and the effectiveness and performance of the proposed networks were compared. The PNN yielded the best classification accuracy.

Chaurasia et al. investigated the prediction of benign and malignant BC by using data mining techniques. They employed three popular data mining algorithms—naive Bayes, RBF network, and J48—to develop prediction models by using breast-cancer Wisconsin having 699 instances, two classes (malignant and benign), including the features of tumor structures and historical data of patients. They reported, according to average accuracy, that naive Bayes exhibits the best capability with 97.36% accuracy; RBF network was second with 96.77% accuracy, and J48 came out third with 93.41% accuracy. Regardless of the various studies that investigated ML for BC prediction, few studies considered the lifestyle and historical data of patients. Few studies considered the combination of the lifestyle, history of diseases, and demographics data to predict BC regarding that these data are more available and affordable. Therefore, in our study, we aimed to predict BC based on the patients’ demographics, history, and lifestyle features by using GA incorporated with several ML approaches such as KNN, RBF, DT, fuzzy neural network (FNN), PNN, and pattern recognition networks. Finally, their performance was compared to introduce the best model for predicting BC.

Materials and Methods

Study design and setting
This retrospective cross-sectional study was conducted in 2021 at Ayatollah Taleghani Hospital, which is the cancer screening hub and treatment center in Abadan city, Southwest region of Khuzestan province, Iran. The study was approved by the Ethics Committee of Abadan University of Medical Sciences (code: IR. ABADANUMS. REC.1400.040). To protect the privacy and confidentiality of patients, we concealed the unique identification information of all patients in the process of data collection. All experiments on the classification
algorithms described in this study were conducted using the Python programming language (version 3.7.7). The Python experiment environment offers a well-defined framework for researchers and developers to run and assess their ML models. The road map of the proposed system for predicting the risk of BC based on lifestyle factors is depicted in Figure 1.

Study participants and sampling

Data preprocessing

It is very important to prepare and clean the dataset before using it to construct ML models. In the preprocessing stage, to ensure effective use of data in classification algorithms, the raw data were inputted using several preprocessing methods such as deletion of missing values, minimum and maximum scalar, and standard scalar. The standard scalar guarantees that every feature has the mean as 0 and variance as 1, bringing all features to the same coefficient. Similarly, in minimum and maximum scalar transfers, the values were such that all attributes are between 0 and 1, and rows with missing values (greater than 50%) were removed. In addition, the remaining missing values were inputted with the mean or mode of each variable. Noisy and abnormal values, errors, duplicates, and meaningless data were checked by researchers in collaboration with two infectious disease specialists and oncologists.

Data sample

A total of 6870 supposed BC cases were referred to this center from February 2017 to 2020 for cancer screening or diagnosis. Of those, 5520 cases underwent mammography. By applying the predefined exclusion criteria, 3930 cases remained. Of those, 1270 patients (32.31%) were introduced as confirmed BC by mammography in combination with biopsy, and 2660 were diagnosed non-BC (67.69%) [Figure 2]. After removing the missing values, two classes (healthy and patient) and 32 integer-valued attributes were identified. The attributes of the missing values, two classes (healthy and patient) and 32 integer-valued attributes were identified. The attributes of the dataset are presented in Table 1.

Feature selection

Feature selection is an effective technique that is used to determine relevant features, reduce the dimensions of the dataset, and improve the efficiency of the classifier. This method, along with the assessment of the technique to score diverse evolved feature subsets, is required to obtain the best or most favorable output.[27] The genetic method, which is a type of feature selection algorithm based on a random optimization technique, was applied to select the most relevant features to predict BC. GA is based on the Darwinian theory, which tries to inspire the strategies of the natural evolution of living beings. The GA operates on a population of solutions at various sequential generations for choosing superior offspring based on the “survival of the fittest” principle. The process begins by making an accidental population of solutions; it does not assess solutions in sequence but assesses a set of solutions synchronously. There are three important operators in GA: selection, crossover, and mutation within chromosomes. In our study, GA
was combined with the best predictive ML algorithms to perform early diagnosis or prediction of BC based on selected features.

Model development and evaluation

Classifiers

In the present study, to predict the risk of BC based on lifestyle factors, several ML classification algorithms, including KNN, RBF, DT, artificial neural network (ANN), FNN, PNN, and pattern recognition, were used. These algorithms were selected because these are simple yet powerful models and can yield feasible results. In addition, the use of different approaches for developing the prediction models may increase the chances of obtaining a better prediction model with high classification accuracy.

ML algorithms usually have a set of parameters that must be set before running the models. The selection of parameters can notably impact the performance of models, but distinguishing the good value of parameters can be complex. In this study, we applied a set of parameters as shown in Table 2.

KNN algorithm

KNN is an algorithm for classifying variables by considering the nearest training data in the feature space. KNN applies an instance-based learning method, which is one of the simplest algorithms among data mining techniques. This method considers the nearest neighbors to each object and decides to dedicate the object to classes.

Artificial neural network

ANN is an ML algorithm that imitates the biological neural network. In our study, we applied two types of networks: multilayer perceptron neural networks (MLPNNs) and RBF networks. The MLPNN maps a set of input data to a set of appropriate output classes. RBF network is another type of neural network. As the input of the neurons in an MLP network takes the weighted sum of its inputs, every input value is multiplied by a coefficient and then the outputs are obtained by summation of the values. A single MLP neuron is a simple linear classifier, but complex nonlinear classifiers can be built by combining these neurons into a network.

Decision trees

DT consists of two parts: nodes and rules. The main idea of this algorithm is to create a tree that contains a root node on top; each non-leaf node represents an attribute, and the final results are represented in the leaf nodes.

Table 1: Data set attributes and corresponding values for BC early detection

| Variable name               | Values       | Variable name               | Values       |
|-----------------------------|--------------|-----------------------------|--------------|
| Age                         | Num          | Meat Consumption Status     | Rarely       |
| Gender                      | Male, Female | Fish Consumption Status     | Rarely, Some times highly |
| Marital Status              | Single, Married | Vitamin Supplements | Yes, No, Some times highly |
| Smoking                     | Yes, No      | Family Breast Cancer History | Yes, No |
| Pregnancy Status            | Yes, No      | Breast Biopsy History       | Yes, No     |
| Having Children             | Yes, No      | Chest Radiology History     | Yes, No     |
| Regular Physical Activity   | Yes, No      | Breast Examination History  | Yes, No     |
| Diabetes                    | Yes, No      | Dairy Products Status       | Rarely, Some times highly |
| Fruit Consumption Status    | Rarely, Some times highly | Colorectal Cancer | Yes, No |
| Having Job                  | Yes, No      | Hyperglyceridemia           | Yes, No     |
| Higher Salt Intake          | Yes, No      | Drinking Alcohol            | Yes, No     |
| Breast Implants             | Yes, No      | Body Mass Index             | Num         |
| Hyperlipidemia              | Yes, No      | Fiber Consumption Status    | Rarely, Some times highly |
| Being overweight or obese   | Yes, No      | Hormone Therapy History     | Yes, No     |
| Vegetable Consumption Status| Rarely, Some times highly | Education Statuses         | High school Bachelor |
| Hypertension                | Yes, No      | Waist                       | Num         |
| Hypercholesterolemia        | Yes, No      | Breast Feeding History      | Yes, No     |
| Contraceptives              | Yes, No      |                             |             |

Figure 2: Flowchart describing patient selection
nodes. DT algorithms have been widely used in data mining applications as they are one of the most powerful classification tools.[33]

Fuzzy neural networks and probabilistic neural networks
FNN integrates the advantages of both fuzzy rule-based systems and neural networks. This hybrid learning algorithm uses a fuzzy inference system and is implemented in the framework of adaptive neural networks. For the classification, for each feature, FNN applies several neurons and membership functions; the number of fuzzy rules is dependent on the number of inputs.[34] PNN is another classification algorithm. This model calculates the distances from the input vector to the training input vectors and generates a vector whose elements indicate how close the input is to a training input for each input in its first layer. The second layer computes the sums of these contributions for each class of inputs and generates as its net output a vector of probabilities. Finally, a complete transfer function on the output of the second layer selects the maximum of these probabilities and outputs 1 for that class and 0 for the other classes.[35]

Evaluation
In this study, we applied 10-fold cross-validation and five performance assessment metrics. Accuracy, specificity, sensitivity, KAPA, error rate, and ROC curve were measured for comparing the performance of the classifiers (Equations 1–4).

To better compare the performance of the algorithms, we assessed the effectiveness of five ML algorithms in terms of time required to build the model, correctly classified instances, incorrectly classified instances, Kappa statistic, mean absolute error, root mean squared error (RMSE), relative absolute error, and root relative squared error (RRSE).

1) classification accuracy = \frac{TP + TN}{TP + TN + FP + FN} * 100

2) classification sensitivity = \frac{Tp}{TP + FN} * 100

3) classification specificity = \frac{TN}{TN + FP} * 100

4) classification error = \frac{FP + FN}{TP + TN + FP + FN} * 100

The output results of the classification accuracy in each run using each of the methods are presented in Table 2. Case class data were considered as negative data, and control class data were considered as positive data. Our dataset contained 32 features.

Ethical consideration
The research deputy of Abadan University of Medical Sciences (ethical code: IR.ABADANUMS.REC.1400.040) approved the current study. To protect the privacy and confidentiality of patients, we concealed the unique identifying information of all patients in the process of data collection and presentation.

Results
Patient selection
We obtained data from 5520 patients in the NCBR. Eighty-eight incomplete case records that had a lot of missing data (more than 70%) were excluded from the analysis. In addition, the missing values were inputted with the mean or mode of each variable. After applying the exclusion criteria, the final analysis was performed on the data of 3930 BC and non-BC cases who were referred to Ayatollah Taleghani Hospital for BC screening and diagnosis. Of the 3390 study participants, 1270 cases (32.32%) were BC patients and 2660 (67.68%) were healthy women, and the median age of participants was 57.25 (interquartile range: 16–86). A flowchart to represent the patient selection methodology is shown in Figure 2.

Result of selected features by genetics algorithm
The modified GA selects the most important and highly related features based on the weights of the features. The experimental results of the modified GA showed that of the 35 included variables, 10 variables were the most significant features in predicting the risk of BC. Table 3 shows the important selected risk factor for BC and their scores.
According to the results, family history of BC, hormone therapy history, breast biopsy history, and dairy product consumption yielded high scores, meaning that these four variables have a high impact on the early prediction of BC.

In this experiment, the selected features of our dataset were checked on a GA-DT modified ML classifier with 10-fold cross-validation methods. The average assessment criteria of 10-fold cross-validation were measured.

As can be seen from Tables 4 and 5, the GA-DT shows good performance, with a mean classification accuracy of 92.9%, a mean specificity of 92.7%, and a mean sensitivity of 91.8%. Table 6 displays the 10-fold cross-validation results of 10 independent runs of the modified GA-DT algorithm with selected features.

**Results of K-fold cross-validation on the dataset with the selected features**

In this study, datasets were used for seven classification models. The dataset was used for training the prediction models, and only 10% of the dataset was for testing. Finally, the mean metrics of 10-fold cross-validation methods were measured. Furthermore, the values of the determining parameters were obtained by running ML classifiers. Table 6 shows the 10-fold cross-validation results and five performance assessment metrics (mean, standard deviation, minimum and maximum classification accuracy, specificity, and sensitivity). The ROC and confusion matrix for the best running of algorithms are presented in Figures 3, 4, and 5.

In this study, for the KNN algorithm, we performed experiments with different values of k (1, 3, and 5). The results revealed that at k = 1, the performance of KNN was superior than that for other values of k, as shown in Table 5. KNN with k = 1 yielded an accuracy of 95.1%, a specificity of 95.9%, and a sensitivity of 90.4. The specificity percent of the KNN algorithm demonstrates the probability that a prediction for BC was negative and the patient does have BC. Furthermore, 90.4% sensitivity demonstrates the probability that the patient was accurately predicted for BC. The ROC and confusion matrix for KNN (k = 1) are depicted in Figure 3.

Table 7 represents the 10-fold cross-validation results and metrics of performance for the GA-DT for the selected feature.

Table 8 shows the classification performance of five other classifiers with 10-fold CV for selected. The confusion matrix and ROC curve of the models are shown in Figures 4 and 5, respectively. The value for the error rate of the classifier and the computation time of classification models on the selected features are shown in Figure 6.

As shown in Table 7 and Figure 5, the performances of all the models are good. The ANN was trained on several inputs and hidden layers and neurons. The ANN model yielded a mean accuracy of 98.6%, a mean specificity of 97.05%, and a mean sensitivity of 99.03% in 10 independent iterations. The standard deviation for the accuracy was 1.1, and the best results in the ten runs were 98.8% accuracy, 97.7% specificity, and 99.1% sensitivity. For the FNN, the mean accuracy in 10 iterations was 97%, with the mean specificity and sensitivity of 98.8%.

### Table 3: Features set selected by Relief-feature selection algorithm and their scores

| Feature name                  | Score |
|-------------------------------|-------|
| Age                           | 0.497 |
| Dairy products status         | 0.515 |
| Family breast cancer history  | 0.610 |
| Breast biopsy history         | 0.531 |
| Chest radiology history       | 0.482 |
| Hormone therapy history       | 0.579 |
| Drinking alcohol              | 0.370 |
| Being overweight or obese     | 0.401 |
| Having children               | 0.468 |
| Education status              | 0.374 |

### Table 4: The performance measures of GA-DT with different parameters in 10 iterations

| Run | Accuracy | Specificity | Sensitivity | Features Selected | The most important feature based on ten independent running of modified GA algorithm |
|-----|----------|-------------|-------------|-------------------|----------------------------------------------------------------------------------|
| 1   | 91.5755  | 87.5312     | 92.7407     | 1,4,5,7,8,9,10,11,12,13,14,17,19,20,21,22,26,28 | Age, dairy products status, family breast cancer history, breast biopsy history, chest radiology history, hormone therapy history, drinking alcohol, being overweight or obese, having children, education statuses |
| 2   | 90.855   | 89.7467     | 91.8519     | 1,4,5,7,10,12,13,14,15,17,18,20,21,25,26,28 | family breast cancer history, hormone therapy history, drinking alcohol, being overweight or obese, having children, education statuses |
| 3   | 90.1062  | 90.1389     | 95.1481     | 1,4,5,7,10,12,17,18,19,20,21,22,25,26,28 | family breast cancer history, hormone therapy history, drinking alcohol, being overweight or obese, having children, education statuses |
| 4   | 90.5049  | 91.6962     | 91.5556     | 1,4,5,7,8,10,11,12,13,14,17,18,19,20,22,24,28,29 | family breast cancer history, hormone therapy history, drinking alcohol, being overweight or obese, having children, education statuses |
| 5   | 92.5788  | 92.3115     | 93.6667     | 1,4,5,7,8,12,14,15,18,20,24,25,27,28,31 | family breast cancer history, hormone therapy history, drinking alcohol, being overweight or obese, having children, education statuses |
| 6   | 90.5788  | 90.4098     | 90.6296     | 1,4,5,7,8,13,14,15,17,18,20,24,25,28,29,31 | family breast cancer history, hormone therapy history, drinking alcohol, being overweight or obese, having children, education statuses |
| 7   | 91.83    | 90.791      | 91.7407     | 1,4,5,7,11,13,14,17,18,20,21,25,27,28 | family breast cancer history, hormone therapy history, drinking alcohol, being overweight or obese, having children, education statuses |
| 8   | 93.2373  | 93.4799     | 93.2593     | 1,4,5,7,9,10,12,14,17,20,21,24,26,28 | family breast cancer history, hormone therapy history, drinking alcohol, being overweight or obese, having children, education statuses |
| 9   | 95.7432  | 95.908      | 94.2407     | 1,4,5,7,8,10,11,14,17,18,20,22,25,28,29 | family breast cancer history, hormone therapy history, drinking alcohol, being overweight or obese, having children, education statuses |
| 10  | 95.5365  | 94.5622     | 94.1667     | 1,4,5,7,11,13,14,17,19,20,22,24,25,27,28,29,31 | family breast cancer history, hormone therapy history, drinking alcohol, being overweight or obese, having children, education statuses |
and 96.9%, respectively. The standard deviation for the accuracy obtained was 0.07, and the best results in the 10 runs were 97.2% accuracy, 99% specificity, and 97% sensitivity. For the pattern net, we obtained the mean accuracy, mean specificity, and mean sensitivity as 98.7%, 97.4%, and 99%, respectively. The standard deviation for the accuracy in 10 iterations was 0.12, and the best results in the 10 runs for the accuracy, specificity, and sensitivity were 98.8%, 97.7%, and 99.1%, respectively. After applying the probabilistic neural network, the mean accuracy, mean specificity, and mean sensitivity were 96%, 91.4%, and 96.9%, respectively. The standard deviation for accuracy was 0.11, and the best measure obtained was 96.2% for the accuracy, and 92.1% and 97% for the specificity and sensitivity, respectively. Table 5 shows that the mean accuracy in 10 iterations of the RBF method was 83%, with the mean specificity and sensitivity of 78% and 84%, respectively. The standard deviation for the accuracy was 1.1, and the best results in the 10 runs yielded an accuracy of 85%. The pattern net model has the best computational time (38 s), and the RBF model has the worst processing time (117 s). The AUC value of the GA-DT model was the highest (99.01) in comparison to other models.

Finally, upon comparing seven ML algorithms, we found that the performance of the GA-DT algorithm was excellent; the second important classification algorithm was pattern net, and the worst performance was noted for RBF.

### System implementation

The system was implemented during February and April 2021. System programming consisted of three types of implementation codes: codes for the user interface implementation, codes for the logic layer implementation, and codes for the database implementation. The user interface in our study comprised four pages: Welcome page (sign up and log-in page), and CDSS module (2 pages). The user interface of the BC risk prediction system was developed using the C# programming language, see Figures 7 and 8.

### Discussion

The prevalence of healthy lifestyle with healthy behaviors is low in all worldwide countries. Thus, the use of an intelligence system for predicting BC risk based on lifestyle factors may assist clinicians in the evaluation of a limited number of important lifestyle features in an endeavor to recognize individuals at high risk for BC. Furthermore, our developed lifestyle-based

| Table 5: Results for running modified GA algorithm in 10 independent executions |
|---|---|---|
| Accuracy | Specificity | Sensitivity |
| Mean | 92.9259 | 92.7407 | 91.88996 |
| Std | 1.806388 | 2.356811 | 1.671002 |
| MIN | 90.6296 | 87.5312 | 90.1062 |
| MAX | 95.7407 | 94.5622 | 94.2407 |

| Table 6: The average performance of 10 independent runs of classifiers based on 10-fold cross-validation |
|---|---|---|---|---|---|
| | K=1 | K=3 | K=5 | K=1 | K=3 | K=5 |
| Accuracy | 8.9668e+01 | 9.3644e+01 | 9.3275e+01 | 9.1157e+01 | 9.4613e+01 | 9.4234e+01 | 9.1935e+01 | 9.5852e+01 | 9.4303e+01 | 8.9350e+01 |
| Specificity | 5.8369e+01 | 1.0816e+01 | 1.0576e+01 | 5.8369e+01 | 1.0816e+01 | 1.0576e+01 | 5.8369e+01 | 1.0816e+01 | 1.0576e+01 |
| Sensitivity | 3.1143e+0-0 | 6.4132e-02 | 8.3126e-02 | 3.0080e-01 | 1.0276e-01 | 9.7362e-02 | 5.8369e-01 | 1.0816e-01 | 1.0576e-01 |
| Std | 8.9160e+01 | 9.3549e+01 | 9.3099e+01 | 9.0671e+01 | 9.4481e+01 | 9.4090e+01 | 8.8693e+01 | 9.5649e+01 | 9.5193e+01 |
| Worst | 9.0304e+01 | 9.3746e+01 | 9.3411e+01 | 9.1677e+01 | 9.4787e+01 | 9.4401e+01 | 9.0472e+01 | 9.5982e+01 | 9.5982e+01 |
| Best | 9.4901e+01 | 9.5852e+01 | 8.9350e+01 | 9.4234e+01 | 9.4613e+01 | 9.4234e+01 | 9.1935e+01 | 9.5852e+01 | 9.4303e+01 |

![Figure 3: The best performance in confusion matrix and ROC curve for GA-KNN algorithm](image-url)
prediction system may enhance consciousness to detect true primordial inhibition through interventions on underlying unwell behaviors to ban the growth of any important risk factors of BC initially rather than treating patients only when the disease advanced.

In the present study, an ML-based predictive system was developed for early risk prediction of BC based on lifestyle factors. The ML algorithms were applied to a preprocessed dataset. Eight well-known classifier algorithms, including KNN (k = 1, 2, and 3), ANN, SVM, FNN, RFB, DT, pattern net, and PNN, were used with a feature selection algorithm (GA) to select the most important predictors. To validate the system, the k-fold cross-validation method was used. To compare the performance of the classifiers, several evaluation metrics derived from the confusion matrix were used. The feature selection method (GA) selects the most important variables that enhance the performance of the classification algorithms in terms of accuracy, specificity, sensitivity, ROC, different metrics of error rate, and the processing time of models.

The most important features of BC were age, consumption of dairy products, BC family history, breast biopsy, chest X-ray, hormone therapy, alcohol consumption, being overweight, having children, and education level. The DT algorithm with 10-fold cross-validation presented the best accuracy of 99.2%, a specificity of 99.5%, and a sensitivity of 97.9% when selected by the GA algorithm. Due to the high performance of the DT algorithm with GA, it was

Table 7: Results for running GA-DT algorithm in 10 independent executions

|               | Accuracy   | Specificity | Sensitivity |
|---------------|------------|-------------|-------------|
| Mean          | 9.9222e+01 | 9.9505e+01  | 9.7949e+01  |
| Std           | 6.4318e-02 | 4.4483e-02  | 1.4923e-01  |
| Worst         | 9.9123e+01 | 9.9426e+01  | 9.7792e+01  |
| Best          | 9.9321e+01 | 9.9554e+01  | 9.8239e+01  |
selected as the core of the clinical decision support system to better predict the risk of BC in terms of accuracy, specificity, and sensitivity.

In terms of specificity, the FNN algorithm has the best specificity among other models with 97% specificity and a standard deviation of 0.07. The best result for sensitivity among the eight models was obtained for pattern net, with a sensitivity of 99.1% and a standard deviation of 0.1. In addition, in terms of computation time, the pattern net with a feature selection (GA) algorithm was the best as compared to the computation time of the other seven algorithms, as shown in Figure 5. Williams et al. [39] showed that data mining approaches have significant predictive power for BC. They indicated that DT had the best accuracy in comparison with other techniques.

In our study, the DT model had a high accuracy of about 99% as well. This may prove the strong power of DT in predicting BC. Another study that compared different data mining algorithms is the research by Higa [40] that selected DT and neural network as the best models for diagnosing benign and malignant tumors of BC with 95% accuracy. Our study also showed that these two approaches (DT and ANN) had the best prediction powers, and we obtained higher maximum accuracy in our study. The proposed ML-GA model in the Jebarani study (2021) showed its effectiveness for classifying benign and malignant BC tumors. [41] Solanki et al. [42] (2021) also conducted a retrospective analysis for BC prognosis by using hybrid supervised ML classifiers.

Finally, the best meaningful results were observed using the J-48 DT-GA classifier for feature selection with an accuracy of 98.83%, MCC = 0.974, sensitivity = 98.95%, specificity = 98.58%, and Kappa statistics = 0.9735.

Several prediction factors have been investigated in previous studies to predict BC, such as breast medical images [43, 44], the biopsy of the lesion [24] and blood tests [45]. However, we considered a more cost-effective approach and available data with the least intervention features for our prediction models. In addition, in a previous study, several unimportant features reduced the accuracy and sensitivity of the prediction system and enhanced the processing time. Therefore, one of the innovations used in this study was applying the feature selection algorithm to select the most important factors that enhance the accuracy, specificity, and sensitivity of the classifiers as well as decrease the running time of the predictive system. In this study, the GA algorithm was utilized for solving these challenges. The results confirm the positive effect of 10 features in predicting the risk of BC, and such satisfactory results can be attributed to the use of GA as a powerful optimizer that selected the best subset features to be included in ML algorithms. It has been inferred that upon hybridizing different ML algorithms, the prediction models show more promising performance compared to a single model. Thus, ML algorithms can be used to construct complex models and make reliable decisions when fed with appropriate features. When there is a valuable set of features, the performance of ML algorithms is anticipated to be adequately acceptable. However, in specific applications, the dataset is often

### Table 8: Classification performance of five other classifiers with 10-fold CV on selected features

| Evaluation criteria | RBF | Probabilistic neural network | Pattern net | FNN | Neural network MLP |
|---------------------|-----|------------------------------|-------------|-----|--------------------|
| Best Time to build a model (s) | 117 | 83 | 38 | 63 | 91 |
| Mean Accuracy (%) | 83.4 | 96.0 | 98.7 | 97.09 | 98.6 |
| Mean Specificity (%) | 77.7 | 91.4 | 97.5 | 98.8 | 97.05 |
| Mean Sensitivity (%) | 84.1 | 96.9 | 99.07 | 96.9 | 99.03 |
| STD | 1.1109e+00 | 1.1832e−01 | 1.2861e−01 | 7.7757e−02 | 1.0787e−02 |
| Worst Accuracy | 81.3 | 95.9 | 98.4 | 97 | 98.5 |
| Best Accuracy | 85.0 | 96.2 | 98.8 | 97.2 | 99.8 |

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**Figure 6:** The error rate of classifiers on the given dataset

**Figure 7:** The welcome page of the clinical decision support system
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Figure 8: The clinical decision support system for predicting the risk of breast cancer

insufficient or imbalanced. Therefore, it is important to train these algorithms and obtain good results with the most relevant set of features.

The predictive BC system identifies persons at high risk based on lifestyle behaviors. For individuals with a high risk of BC, protocols are in place for the optimal period of treatment. However, for persons with a low risk of BC, our developed system can provide information about long-term BC risk and BC overall burden. Ultimately, future studies are needed to evaluate the feasibility and impacts of our lifestyle BC risk prediction system, BC risk feature improvement, and overall BC risk evaluation when integrated into the clinical care environment, particularly in integration with other clinical-based risk systems.

Limitations and reconsecrations
This study had some limitations that are necessary to be recognized. Dealing with a retrospective-single center dataset, the present study suffered from the low quality (imbalanced, noisy, duplicates, and meaningless values), low quantity (missing cells), and non-optimal generalizability of data in the selected database. First, we removed noises, duplicates, and meaningless records manually as much as possible from the dataset. To solve the imbalanced dataset problem, by using the SMOTE technique, the bias was minimized via class balancing. Second, missing values were imputed with the mean or mode of each variable. Finally, it is recommended to use a dataset with a larger sample size in a multi-center setting in future studies.

Despite the limitations of our study, this is an important study on clinical prediction systems assessing the incidence risk of BC by routine lifestyle features in Iran. We expect this prediction clinical decision support system to play a significant role in enhancing the quality of decision-making and detecting individuals at high risk for BC and applying BC prevention strategies in the field of health care policy in our countries, where the screening program at the early stages is not included in the routine national health program. Additionally, the main novelty of our study is that we derived the lifestyle-based BC risk prediction empirically by using state-of-the-art and novel hybrid ML methods (i.e., hybrid GA-DT) by considering various features simultaneously. Moreover, we considered a Windows-based application to enable BC risk prediction in the first line of the healthcare system.

Conclusion
The predictive BC system identifies persons at high and elevated risk for BC based on lifestyle behaviors and can be used as an essential clinical screening tool for the early prevention of BC and serve as important tools for developing preventive health strategies. However, there is also an essential need to perform studies to evaluate the feasibility and impacts of such a lifestyle-BC risk prediction system, especially when integrated into the clinical care environment, particularly in integration with other clinical-based risk systems.

Ethics approval and consent to participate
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Consent for publication
Not applicable.

Availability of data and materials
All data generated and analyzed during the current study are not publicly available but are available from the corresponding author on reasonable request and the Research Committee of Abadan University of Medical Science’s approval.

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Abbreviations
ML: Machine Learning, GA: Genetic Algorithm, BC: Breast Cancer, ROC: Receiver Operating Characteristic, WHO: World Health Organization, LR: Logistic Regression, SVM: Support Vector Machine, KNN: K-Nearest Neighbors, DT: Decision Tree, RF: Random Forest, AdaBoost: Adaptive Boosting, GBM: Gradient Boosting Machine, XGBoost: eXtreme Gradient Boosting, SOM: Self-Organizing Map,
RBF: Radial Basis Function Network, GRNN: General Regression Neural Network, and PNN: Probabilistic Neural Network, WBCD: Wisconsin Breast Cancer, FNN: Fuzzy Neural Networks, ANN: Artificial Neural Network, RMSE: Root Mean Squared Error, RRSE: Relative Absolute Error, and Root Relative Squared Error.

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Conflicts of interest
There are no conflicts of interest.

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