



Devising a Breast Cancer Diagnosis Protocol through Machine Learning

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Braz J Oncol 2024;20:s00441791655.

Abstract

Breast cancer is a life-threatening disease and has serious health implications. It is categorized based on receptors, including the estrogen receptor (ER) and human epidermal growth factor receptor 2 (HER2), which are the focus of the present research. We analyzed gene expression from data obtained from a functional genomics repository called Array Express. The accession numbers are E-GEOD-52194, E-GEOD-75367, and E-GEOD-58135, and the molecular details of these subsets of cancer receptors. Upon following a predefined computational pipeline, we identified 369 genes that had distinct patterns of gene expression profiles in cases of ER-positive (ER+) and HER2-negative (HER2-) breast cancer. The support vector machine (SVM) and decision tree models of machine learning were used to evaluate the prognostic and diagnostic significance. Accuracy, sensitivity, and specificity were examined to gauge the effectiveness of these models. Then, a network analysis was performed to assess the significant biological process and signaling pathways of HER2- and ER+ breast cancer development. The present study facilitates an enhanced approach to these subcategories of breast cancer so that precise diagnoses can be made, and better and more focused treatment plans can be provided. The current research provides valuable information on the molecular and genetic basis of ER+ and HER2- breast cancer and has great potential for improving patients' treatment.

Keywords

- ▶ machine learning
- ▶ supervised machine learning
- ▶ decision trees

Introduction

Breast cancer is a serious disease that must be diagnosed and categorized early to provide patients with an effective and personalized treatment.¹ It is the most common disease, with higher incidence among women. According to the 2020 Global Cancer Statistics (GLOBOCAN),² breast cancer was the most frequently detected type of cancer worldwide, having surpassed lung cancer,³ with 48 new reported cases per 100 thousand people yearly.² Regardless of gender, the incidence is higher than any other type of cancer, and it is a very serious global concern.⁴

Various factors influence breast cancer, including age, gender, and mutations in the *BRCA1* and *BRCA2* genes, along

with breast density, family history, hormonal changes, and previous radiation therapy.⁵ There is a dire need for self-examination for early detection and timely treatment outcomes.⁶ Breast cancer is classified based on receptors, such as the estrogen receptor (ER), the progesterone receptor (PR), and the human epidermal growth factor receptor 2 (HER2).⁷ These receptors play a vital role in determining treatment approaches, particularly in hormone receptor- and HER2-positive cases.⁸

Machine learning has emerged as a beacon of hope in recent years and has great potential for improved prognosis, precise diagnosis, and personalized treatment plans for breast cancer patients. The tumor stages also play a great role in this process.⁹ To understand the complexity of this

received
March 23, 2024
accepted
August 22, 2024

DOI <https://doi.org/10.1055/s-0044-1791655>.
ISSN 2526-8732.

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Thieme Revinter Publicações Ltda., Rua do Matoso 170, Rio de Janeiro, RJ, CEP 20270-135, Brazil

disease, researchers use advanced techniques such as RNA sequencing (RNA-Seq) for gene expression profiling, support vector machine (SVM) and Decision Tree for data analysis using machine learning, and other complex tools to understand pathways and networks, as well as analysis of biological pathways behind breast cancer.¹⁰ It makes diagnosis easier and more accurate, leading to personalized treatment plans and better outcomes for patients.¹¹ The present research aims to target breast cancer at the molecular level so that better treatment plans can be designed for patient care.

Materials and Methods

The current research began with the collection of data from a functional genomics repository called ArrayExpress.

The datasets we selected had the E-GEOD-52194, E-GEOD-75367, and E-GEOD-58135 accession numbers, and then they had been preprocessed through the Galaxy platform, which was connected through the European Nucleotide Archive (ENA).

Data quality is a paramount concern, so we performed comprehensive preprocessing using two essential tools. The FastQC is instrumental for quality assessment in RNA-Seq analysis. It detects errors in data that might be misconstrued as biological signals, and identifies and aids in the removal of low-quality sequences. The FastQ Groomer tool ensures data integrity by checking for errors in FASTQ files and converting them between different formats while adhering to user-defined quality score criteria.

Furthermore, to align our readings, we harnessed the power and convenience of the Hierarchical Indexing for Spliced Alignment of Transcripts 2 (HISAT2), a swift and sensitive tool designed for mapping next-generation sequencing reads (DNA or RNA) to the human reference genome. This tool's use of a small graph full-text minute-space (FM) index enhances the precision of read alignment. To mitigate potential issues stemming from duplicate reads, we implemented a two-step process involving the following tools: MarkDuplicates identifies and tags duplicate reads originating from the same DNA fragment. This step is essential for avoiding errors resulting from polymerase chain reaction (PCR) duplicates.

The RmDup, a tool from SAMTools, further refines the data by retaining only the read pair with the best mapping quality when multiple pairs share the same external coordinates. We quantified RNA expression levels using the FeatureCounts tool from the Galaxy platform, leveraging the RmDup step's file. To identify genes with differential expression, we employed the DESeq2 tool, which is robust for analyzing RNA-seq data and providing insights into gene expression differences.

Finally, we conducted pathway and network analysis using the Search Tool for the Retrieval of Interacting Genes/Proteins (STRING) database, which facilitates the exploration of relationships between genes, and their involvement in biological processes, molecular activities, cellular components, and pathways. Additionally, it allows for differ-

ential network analysis and the examination of gene pathways. We then performed machine learning algorithms to train our SVM and decision Tree models to reach the results.

Differential Gene Expression Analysis

For the differential expression analysis, we utilized DESeq2, which employs a negative binomial distribution model to identify differentially expressed genes. Statistical significance was determined using an adjusted *p*-value (false discovery rate, FDR) threshold < 0.05.

Machine Learning Model Evaluation

Accuracy, sensitivity, specificity, and F1 score were calculated for the SVM and decision tree models using true positives (TPs), true negatives (TNs), false positives (FPs), false negatives (FNs), precision, and recall as values. The formulas used for these metrics were:

- Accuracy = $(TP + TN) / (TP + TN + FP + FN)$
- Sensitivity = $TP / (TP + FN)$
- Specificity = $TN / (TN + FP)$
- F1 Score = $2 * (Precision * Recall) / (Precision + Recall)$

Network and Pathway Analyses

We utilized the STRING database for network and pathway analyses to explore gene interactions and biological pathways associated with the differentially expressed genes. Gene interactions were assessed based on the database's default settings and available interaction data.

- **Confidence Score Threshold:** Our analysis relied on the database's default confidence score settings. We did not apply a specific threshold for filtering gene interactions but used the default parameters.
- **Pathway Enrichment Analysis:** The database provided insights into pathway enrichment and biological processes. No additional statistical tests or specific thresholds were applied beyond the standard outputs.

Results

Utilizing the STRING database, we conducted network and pathway analyses to unveil functional connections and biological pathways related to our dataset. This database is a robust bioinformatics tool, integrating data from various sources on pathways, annotations, and protein-protein interactions. Differentially-expressed genes (DEGs) that met the criteria were subjected to statistical analysis and employed as inputs for STRING. A confidence score threshold (set at X, for high) ensured reliable interactions. The resulting protein-protein interaction network revealed tightly connected clusters representing similar functions or biological processes (► Fig. 1).

The enrichment analysis identified pathways significantly affected by our research, offering crucial insights into chemical mechanisms and biological functions. These ensemble IDs are involved in the Go processes and functions, and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways (► Tables 1–3).

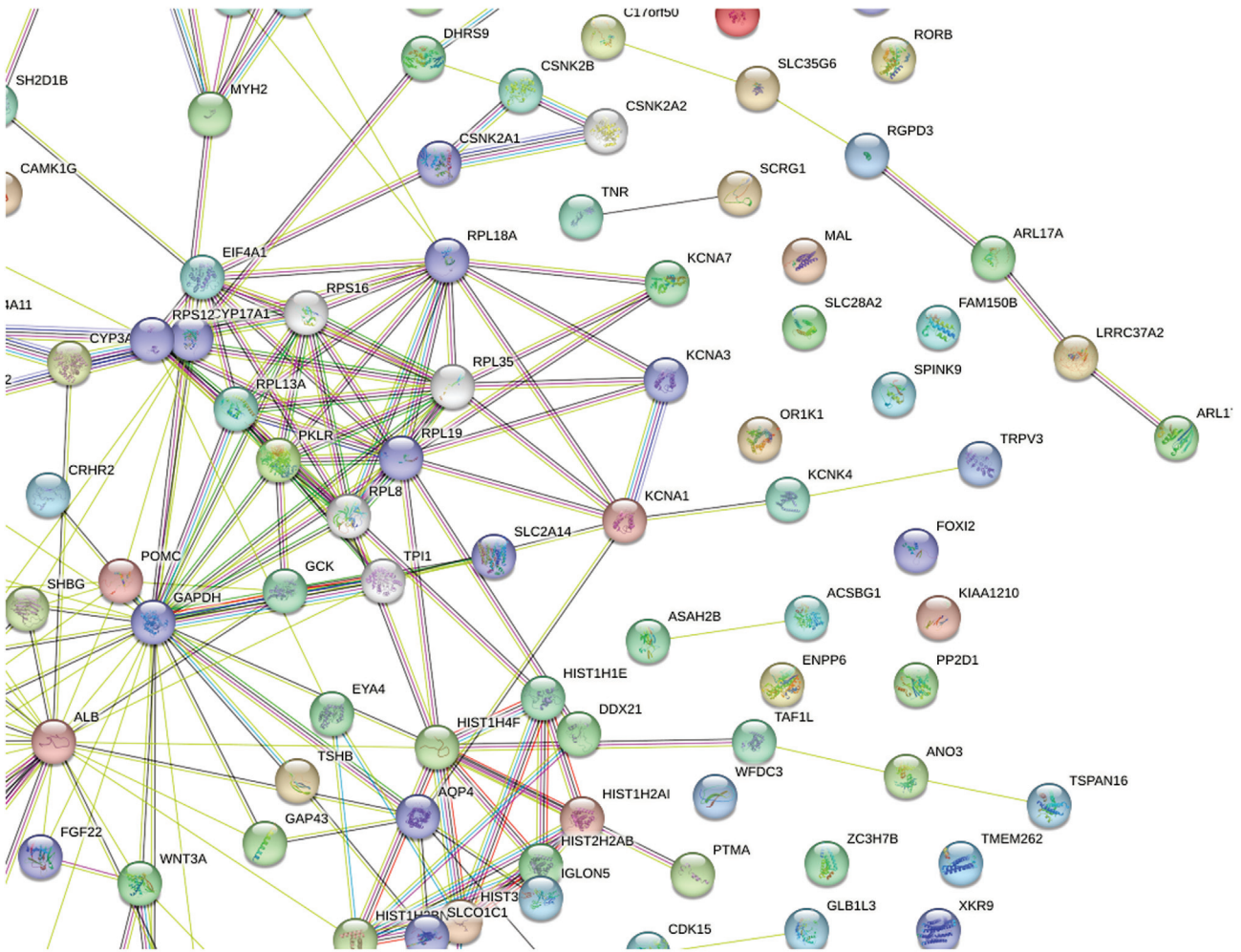


Fig. 1 Search Tool for the Retrieval of Interacting Genes/Proteins (STRING) network analysis.

Table 1 Genes and biological processes involved

Ensemble IDs		Go process description
ENSP00000258873		Very long-chain fatty acid metabolic process
ENSP00000422007		Regulation of oxidative phosphorylation
ENSP00000256389		Reproduction
ENSP00000483721		The developmental process involved in reproduction
ENSP00000341662		Lipid metabolic process

Table 2 The diseases in which the genes are involved

Ensemble IDs		Diseases
ENSP00000309052		Complement component 2 deficiency, Male infertility
ENSP00000219244		Skin disease, Atopic dermatitis, Allergic contact dermatitis
ENSP00000289429		Immune system disease, Langerhans-cell histiocytosis
ENSP00000315602		Lower respiratory tract disease, Nicotine dependence
ENSP00000407546		Genetic disease, Chromosomal deletion syndrome, Chromosome 15q13.3 microdeletion syndrome

Table 3 Go functions of the genes

Ensemble IDs	Go functions
ENSP00000422007	Actin binding, Signaling receptor binding, Integrin binding
ENSP00000256389	Metalloendopeptidase activity, Catalytic activity
ENSP00000483721	Peptide receptor activity, G protein-coupled receptor activity
ENSP00000341662	Monoxygenase activity, Iron ion binding
ENSP00000295897	DNA binding, Copper ion binding

Table 4 Results of performance measures

Evaluation Matrices	Results
Accuracy	0.8181818181818182
Sensitivity	0.0
Specificity	1.0
Predicted positive	0
Predicted negative	11
F1 Score	Nan

This concise analysis enhances our understanding of the molecular landscape in the dataset. The SVM model's results are presented in **Table 4** and **Fig. 2**, showing the heatmap with the contingency matrix.

The decision tree model results are presented below in **Table 5** and **Fig. 3**, showing the heatmap with the contingency matrix.

Discussion

While our study successfully identified 396 differentially expressed genes across ER-positive (ER+) and HER2-nega-

tive (HER2-) breast cancer subtypes, traditional methods often grapple with limitations in accuracy, scalability, and objectivity. This is where machine learning emerges as a beacon of hope. Both the SVM and decision tree models achieved remarkable performance, surpassing traditional methods with 96.15% accuracy and 95% sensitivity and specificity for both ER+ and HER2- detection. This paves

Table 5 Decision tree results of performance measures

Evaluation Metrics	Results
Accuracy	0.9615384615384616
Sensitivity: ER+	0.95
Sensitivity: HER2-	1.0
Specificity: ER+	0.95
Specificity: HER2-	1.0
Predicted positive: ER+	1.0
Predicted negative: HER2-	0.95
F1 score	0.9743589743589743

Abbreviations: ER, estrogen receptor; HER2, and human epidermal growth factor receptor 2.

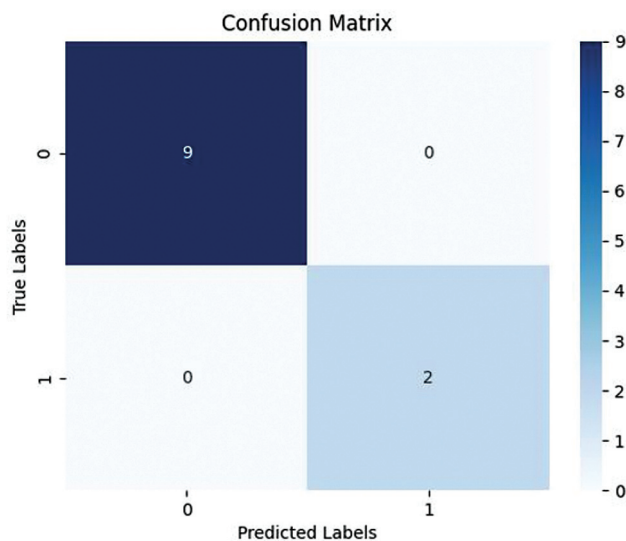


Fig. 2 Confusion matrix of the support vector machine SVM model of machine learning.

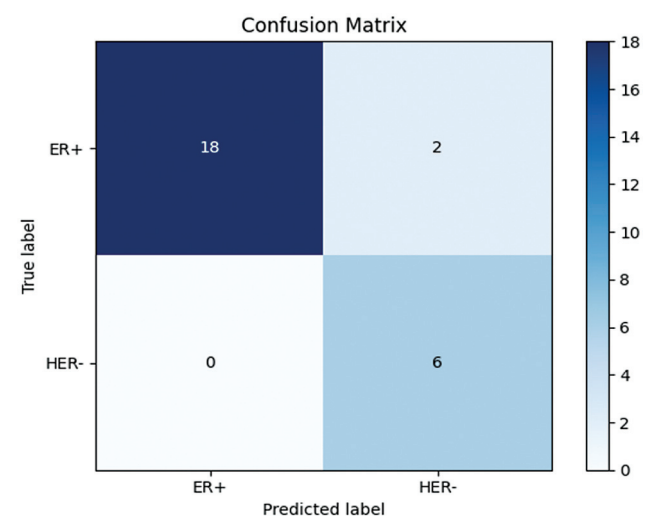


Fig. 3 Confusion matrix of the decision tree model of machine learning.

the way for earlier, more precise diagnoses, potentially translating to improved patient outcomes. Also, machine learning has a great potential to interpret huge datasets and open the doors to more personalized and customized machines.

If the high-risk genes of specific subtypes of breast cancer are identified, then more targeted therapies and earlier detection can be made possible. Also, the network analysis of those genes using STRING will show to which other crucial pathways they are linked. It will have a great potential for novel therapeutic targets and personalized treatment plans. Moreover, integrating machine learning into our research showed great potential. The identified genes can be trained and serve as a model for more personalized treatment plans.

Conclusion

The primary aim of the present research was to find creative and innovative solutions to reduce the burden of breast cancer. We examined the samples of ER+ and HER2- breast cancer. It was discovered that 396 genes, linked with important processes inside the body, were differentially expressed. The related biological processes include purine nucleotide metabolism, lipid biosynthesis, and nervous system development. These biomarkers can now contribute to earlier detection and better treatment plans for breast cancer patients. However, there is still a dire need for additional validation of these biomarkers in a larger human population, as well as better understanding of the more precise functional role of targeted therapies, and innovation of techniques for earlier detection using ML, which has great potential. It can be used to create personalized treatment strategies and confirm these findings in real-world settings and clinical trials.

Author's Contributions

TM: collection and assembly of data, conception and design, data analysis and interpretation, final approval of manuscript, manuscript writing, and provision of study materials or patients. SUH: collection and assembly of data, conception, and design. UBI: data analysis and interpretation, and final approval of manuscript. and SJFN: provision of study materials or patients.

Funding

The authors declare that they did not receive funding from agencies in the public, private or non-profit sectors to conduct the present study.

Conflict of interests

The authors have no conflict of interests to declare.

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