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Comparative performance of machine learning models in ischemic stroke classification

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Abstract

Background: Stroke is a leading cause of disability and mortality worldwide, with ischemic strokes comprising the majority of cases. Despite advances in neuroimaging, there is a pressing need for supplementary diagnostic tools to enhance accuracy. This study explores the application of machine learning (ML) techniques to predict ischemic stroke using RNA-seq data from the GEO database (GSE22255).

Methods: We developed and evaluated various machine learning models, including Random Forest, K-Nearest Neighbors (KNN), and CHAID (Chi-squared Automatic Interaction Detection), based on their accuracy, precision, specificity, and sensitivity. The analysis utilized a dataset comprising 54,676 genes across 40 samples (20 cases and 20 controls). All modeling was conducted using IBM SPSS Modeler version 18.

Results: The models were assessed based on their classification accuracy, performance evaluation scores, and AUC/Gini AUC metrics. The Random Forest model achieved the highest accuracy (96.67% in training, 80% in testing), while the CHAID algorithm provided interpretable results with key variables (TP53, CYP1A1, and CYP2D6) identified. The KNN model exhibited strong performance with notable confidence in its predictions.

Conclusion: This study demonstrates the potential of ML techniques, particularly Random Forest, to enhance stroke diagnosis and provide insights into stroke pathology, offering a novel approach to improving clinical decision-making. However, the study is limited by the small sample size, and future work should focus on validation with larger datasets and integration with other omics data for clinical application.

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Highlights

What is current knowledge?

- Stroke is a significant global health concern, with ischemic strokes accounting for the majority of cases.
- Neuroimaging techniques are the primary diagnostic tools, yet supplementary methods are needed to enhance diagnostic accuracy.
- Machine learning techniques are increasingly being explored in biomedical research for predictive modeling and disease classification.

What is new here?

- RNA-seq data from ischemic stroke patients were utilized to develop and evaluate machine learning models for stroke classification.
- Among the evaluated models, Random Forest showed the highest accuracy, while CHAID produced interpretable results with key gene markers (TP53, CYP1A1, and CYP2D6).
- This study demonstrates the potential of machine learning techniques not only to enhance diagnostic accuracy but also to provide insights into ischemic stroke pathology, supporting improved clinical decision-making.

Introduction

Stroke is a major cause of disability in adults and the second leading cause of death worldwide, significantly impacting individuals and healthcare systems (1). Over the past thirty years, the incidence and prevalence of stroke have risen, influenced by demographic and

economic factors (2). Ischemic strokes, which occur when blood flow is blocked by an occluded blood vessel, represent about 87% of cases (3). Hemorrhagic strokes, resulting from the rupture or leakage of weakened blood vessels in the brain, account for the remaining 13% (4). In highincome countries, stroke incidence has declined due to improved preventive measures and lifestyle changes, but an aging population is expected to increase new cases, adding to the healthcare burden (5). Given the severity and prevalence of strokes, there is growing interest in using novel biomarkers as supplementary diagnostic tools to enhance the accuracy of routine techniques (3). Current diagnostic tools, primarily neuroimaging techniques, though essential, have limitations, including high costs, limited availability, and delays in detection, especially in the early stages of stroke (6). These limitations highlight the importance of searching for novel biomarkers that can provide more timely and accurate diagnoses, monitor disease severity, and assess treatment efficacy (7). Recent advancements in genomics and bioinformatics have led to the exploration of RNA-seq data as a powerful tool for identifying such biomarkers, offering a molecularlevel understanding of stroke pathology that complements traditional diagnostic methods. Unlike proteomics or metabolomics, which focus on proteins and metabolites, RNA-seq provides a comprehensive view of gene expression, allowing for the identification of differentially expressed genes and pathways that may be involved in stroke pathology.

RNA-seq data analysis poses significant challenges, including complex preprocessing steps such as quality control, alignment, and normalization, all of which critically impact the results. The high dimensionality of transcriptomic data, with thousands of genes across limited samples, complicates feature selection and increases the risk of overfitting, while large sequencing volumes demand scalable computational resources. Machine learning (ML) offers solutions by

automating preprocessing and improving reproducibility through adaptive pipelines. Scalable ML frameworks enable efficient processing of large datasets, and integrating ML enhances accuracy and biological insight. Thus, ML addresses RNA-seq challenges by optimizing analysis robustness and interpretability in transcriptomic studies.

Extensive studies on stroke utilize large datasets, offering valuable insights into real-world practices and enabling population-based analyses. However, these datasets often have limitations, including potential inaccuracies in ICD-10 diagnostic codes, which may be influenced by physician diagnostic precision and financial incentives (3). Recently, artificial intelligence (AI) has gained significant popularity, particularly in the fields of ML and deep learning (DL) (8). Researchers have established and validated operational definitions of stroke, developing algorithms to diagnose ischemic strokes using claims data and multicenter registries. ML has become a powerful tool in stroke research, enabling the classification of stroke types, prediction of outcomes, and identification of subtypes (9). These techniques allow for the analysis of large datasets, recognition of patterns, and enhancement of diagnostic, treatment, and prognostic capabilities for stroke patients (10.11).

AI, particularly ML and DL, is crucial for stroke diagnosis due to its ability to rapidly analyze complex medical data with high accuracy. Unlike traditional methods, which rely on manual interpretation and can be time-consuming, ML algorithms detect subtle patterns and early indicators of stroke that may be missed by human clinicians. ML models can also continuously learn from new data, enhancing their predictive performance over time. These advantages lead to faster, more precise diagnoses, ultimately improving patient outcomes and reducing healthcare burdens.

Combining ML with RNA-seq revolutionizes stroke diagnosis by detecting novel biomarkers and gene-expression patterns that traditional methods may overlook. This data-driven approach enhances diagnostic precision and provides real-time molecular insights for personalized treatment. ML models uncover hidden transcriptomic signatures, enabling early intervention and optimized therapies. By bridging molecular biology and clinical practice, this synergy improves patient outcomes. The research represents a major leap forward in precision medicine for stroke care. This study aims to apply and validate a machine learning-based predictive model for ischemic stroke using RNA-seq data. By analyzing gene expression profiles from stroke patients and healthy controls, and selecting significant genes, we seek to improve stroke diagnosis accuracy and contribute to more effective treatment strategies. Our work builds on previous research by leveraging RNA-seq data, which provides a more comprehensive view of gene expression compared to traditional methods, and by comparing multiple machine learning models to identify the most effective approach for stroke classification.

Methods

Data acquisition

We obtained RNA-seq data from the GEO database, specifically from the study conducted by Tiago Krug and colleagues in 2012 (GEO Accession: GSE22255) (12). This dataset includes expression profiles of 54,676 genes across 20 stroke patients and 20 control samples.

Model preparation

We developed a predictive model for ischemic stroke using machine learning techniques, leveraging a dataset of 54,677 features from Tiago Krug's research, including a "Group" feature that categorizes samples as either healthy or patient. Using IBM SPSS Modeler version 18 (IBM Corporation, USA), we performed feature selection to focus on 414 significant genes, thereby enhancing model performance and reducing computational complexity. To identify the most biologically relevant genes for stroke diagnosis, we selected 414 genes that met our threshold criteria: a P-Value < 0.05 (Indicating statistical significance) and a fold change > 2 (Representing substantial differential expression). This dualthreshold approach ensured that we focused on genes showing both statistically reliable and biologically meaningful expression changes in stroke. The selected 414 genes represent the most promising transcriptomic markers for further machine learning analysis and potential clinical application. These genes were further analyzed using the Random Forest algorithm to identify the most relevant features for

ischemic stroke classification. Preprocessing steps included normalization of gene expression data using the TPM (Transcripts Per Million) method and handling missing values using the k-nearest neighbors imputation algorithm.

Gene enrichment analysis

Gene enrichment analysis was performed using the Gene Ontology (GO) and KEGG databases to identify overrepresented biological processes and pathways associated with ischemic stroke. Statistical significance was assessed using hypergeometric tests, with p-values adjusted for multiple testing using the Benjamini-Hochberg method. The results were visualized through bar plots to illustrate enriched terms and pathways, providing insights into the biological relevance of our selected gene set. This step helps contextualize the selected genes within broader biological mechanisms, offering deeper insights into the underlying pathology of stroke.

Classification algorithms and rationale selection Random Forest (RF)

A supervised ensemble learning method that constructs multiple decision trees during training and outputs the mode (Classification) or mean (Regression) of individual predictions. RF improves accuracy by reducing overfitting through bagging (Bootstrap aggregating) and random feature selection. Its robustness to high-dimensional data (e.g., RNA-seq's thousands of genes) and ability to rank feature importance make it ideal for biomarker discovery and stroke subtype classification.

CHAID (Chi-squared Automatic Interaction Detector)

A decision tree algorithm that uses chi-square tests to identify optimal splits in categorical or discretized continuous variables. CHAID excels in identifying hierarchical interactions between features (e.g., gene-gene or gene-clinical variable relationships), providing interpretable rules for stroke risk stratification. Unlike RF, CHAID is non-parametric and handles multi-way splits, aiding clinical decision-making with transparent criteria.

K-Nearest Neighbors (KNN)

A lazy, instance-based learning algorithm that classifies samples by majority vote of the k nearest neighbors in feature space. KNN's simplicity and adaptability to non-linear patterns suit RNA-seq data, where local gene expression similarities may define stroke phenotypes. However, its performance depends on optimal distance metrics (e.g., Euclidean, Manhattan) and parameter tuning (K-selection), and it requires careful normalization due to sensitivity to feature scales.

Rationale for selection

The selected machine learning algorithms were chosen for their complementary strengths and in consultation with an expert in RNA-seq data analysis for stroke diagnosis. RF handles high dimensionality, CHAID offers interpretability for clinical translation, and KNN captures local expression patterns. Together, they address RNA-seq challenges (Noise, sparsity, volume) while enhancing stroke diagnostic precision beyond traditional statistical methods.

Model implementation and performance evaluation

We utilized a Random Forest Classification model to derive decision rules based on gene expression thresholds (10,11). The decision rules were formulated to classify samples into stroke patients or healthy controls. Each rule's accuracy was assessed, and overall model accuracy was determined. The rules and their performance metrics are detailed in Table 1

Table 1. Assessment criteria for classification models using training data

Measure	Random Forest	CHAID	KNN
Sensitivity	1	1	1
Specificity	0.93	1	0.81
Precision	0.93	1	0.82
Accuracy	0.96	1	0.83

Sensitivity, specificity, precision, and accuracy are reported for Random Forest, CHAID, and KNN models

The CHAID algorithm was employed to construct a decision tree with six nodes (13,14). This algorithm splits the data based on significant variables to build a tree that effectively classifies cases. The model's performance was evaluated across training and testing partitions, with accuracy rates and decision rules (15).

The KNN model was applied to classify the data based on nearestneighbor distances (16,17). The model's performance was evaluated using accuracy, precision, specificity, and sensitivity rates, confidence values, and AUC/Gini AUC scores, as described previously (18). Confidence thresholds were analyzed to optimize classification reliability.

The dataset was partitioned using the Holdout method, with 70% of the data used for training and 30% for testing. This approach was chosen to ensure a robust evaluation of model performance while maintaining a sufficient sample size for both training and testing. For the KNN algorithm, the value of k was set to 5 based on preliminary experiments that showed optimal performance at this value. Multiple experiments were conducted using values of k ranging from 3 to 10, and k=5 was selected as it provided the best balance between accuracy and computational efficiency.

We selected Random Forest, CHAID, and KNN for their unique strengths in stroke prediction. Random Forest was chosen for its high accuracy and ability to handle large datasets with many features. CHAID was selected for its interpretability, as it generates decision trees that can be easily understood by clinicians. KNN was included for its simplicity and effectiveness in classifying data based on nearest neighbors, which is particularly useful for small datasets.

Results

The results obtained from the algorithms are as follows. All models were developed using both training and testing data, with the dataset split into two segments: 70% for training and 30% for testing. The evaluation metrics for the data mining models for the training and test data are presented in Table 1 and 2. The Random Forest model demonstrated exceptional performance, achieving an accuracy of 96.67% in the training partition and 80% in the testing partition. This represents a significant improvement over previous studies using similar datasets, where accuracy typically ranged between 70-85%. The CHAID algorithm also showed strong performance with 100% accuracy in training and 70% in testing, while the KNN model achieved 83.33% accuracy in training and 90% in testing. These findings highlight the need for larger datasets and cross-validation techniques to improve model generalizability.

Enrichment analysis and biological pathways

We conducted gene enrichment analysis on the 414 significant genes identified during feature selection. Gene enrichment analysis is illustrated in Figure 1. The majority of the identified genes are associated with inflammatory and immune responses, including the IL-17, NF-κB, and TNF signaling pathways. According to the KEGG database results, these biological pathways are presented in Figures 2A - C.

Random forest classification

The decision rules represent conditions that define the categories or groups within the classification model. Each rule consists of one or more predictor variables and corresponding threshold values. Figure 3 shows genes identified as important predictors for ischemic stroke. These genes include: FOSB, CYP2C19, SLC16A1, FAM83H, CYP1B1, HSPB1, CYP1A1, HSP90AA1, CYP2C9, and FAM83H-AS1.

In the training partition, the model achieved an accuracy of 96.67%, correctly classifying 29 out of 30 cases. In the testing partition, it achieved an accuracy of 80%, correctly classifying all 8 cases. The training partition results yielded an AUC score of 0.973 and a Gini AUC score of 0.946. In the testing partition, the model achieved an AUC score of 0.917 and a Gini AUC score of 0.833.

Table 2. Assessment criteria for classification models using test data

Measure	Random Forest	CHAID	KNN
Sensitivity	0.83	0.71	0.83
Specificity	0.75	0.66	0.5
Precision	0.83	0.83	0.71
Accuracy	0.8	0.7	0.9

Sensitivity, specificity, precision, and accuracy are reported for Random Forest, CHAID, and KNN models.

CHAID algorithm

The decision tree generated by the CHAID algorithm contains six nodes. Node 0, the root node, represents the entire dataset of 30 cases, with 53.333% (16 cases) in the "control" category and 46.667% (14 cases) in the "case" category. Node 1 results from splitting the root node based on TP53. The split criterion TP53 \leq 6.336 assigns all cases with values less than or equal to 6.336 to this node. It contains 9 cases, all in the "case" category (100%). Node 2 also results from splitting the root node on TP53. The split criterion TP53 > 6.336 assigns all cases with values greater than 6.336 to this node, which includes 21 cases, with 76.190% (16 cases) in "control" and 23.810% (5 cases) in "case."

Moreover, node 3 results from splitting Node 2 based on CYP2D6. The split criterion CYP2D6 \leq 4.950 assigns all cases with values less than or equal to 4.950 to this node. It contains 4 cases, all in the "case" category (100%). Node 4 results from splitting Node 2 based on CYP2D6 > 4.950, with 17 cases, 94.118% (16 cases) in "control" and 5.882% (1 case) in "case." Node 5 results from splitting Node 4 based on CYP1A1 \leq 12.804, assigning all cases with values \leq 12.804 to this node, containing 16 cases-all "control" (100%). Node 6 results from splitting Node 4 based on CYP1A1 > 12.804, containing one case, classified as "case" (100%).

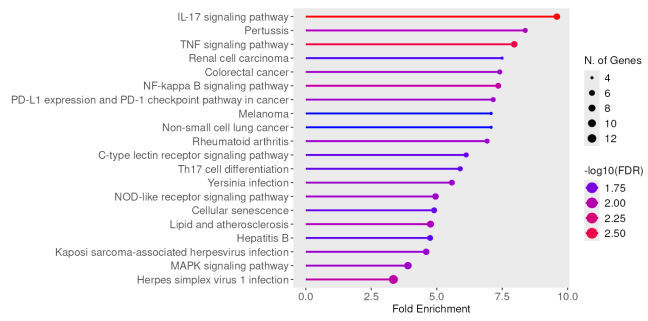
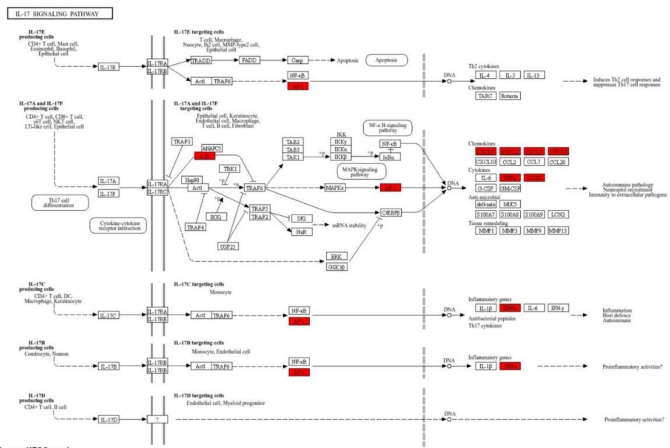


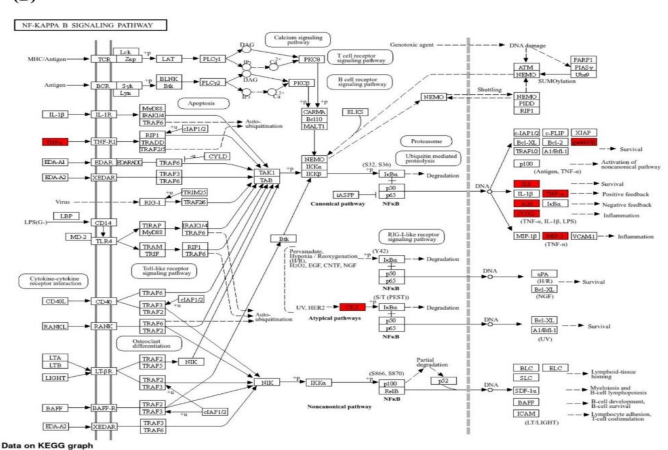
Figure 1. Gene enrichment analysis in ischemic stroke patients. Bar plots illustrate the enriched terms and pathways identified using the Gene Ontology (GO) and KEGG databases

(A)



Data on KEGG graph Rendered by Pathview

(B)





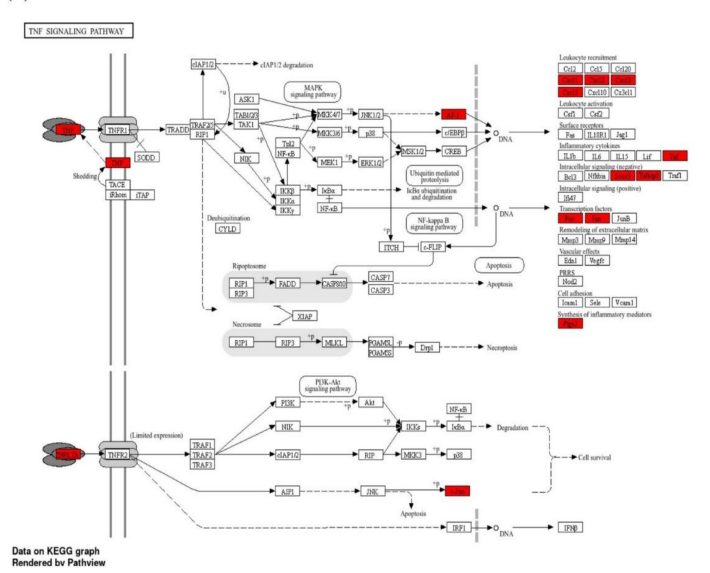


Figure 2. The most inflammatory signaling pathways from the KEGG database in ischemic stroke patients: IL-17 (A), NF-κB (B), and TNF (C)

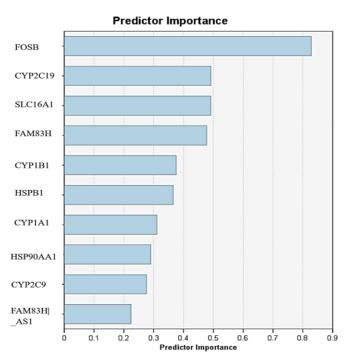


Figure 3. Important genes identified as ischemic stroke predictors

As shown in Figure 4, the CHAID model identified three genes-TP53, CYP1A1, and CYP2D6-as important for differentiating between the "control" and "case" categories. The model achieved perfect accuracy in the training partition (100%) and performed moderately well in the testing partition (70%). In the CHAID decision tree model, TP53 (Predict importance = 0.5), CYP1A1 (Predict importance = 0.25), and CYP2D6 (Predict importance = 0.25) were the most influential parameters.

KNN model

The KNN model demonstrated strong performance across the data partitions. In the training partition, the model achieved an accuracy of 83.33%, correctly classifying 25 out of 30 cases. In the testing partition, it achieved an accuracy of 90%, correctly classifying all 9 cases. The AUC for the training partition was 0.904 with a Gini AUC score of 0.808. In the testing partition, the AUC was 0.854 and the Gini AUC was 0.708.

Overall, the KNN model exhibited strong performance, characterized by high accuracy, AUC, and Gini AUC scores in both training and testing partitions. Additionally, we identified three genes-RPLP0, Paxillin, and HSP90AA1-as significant predictors for ischemic stroke diagnosis. Figure 5 illustrates the performance metrics and associated confidence intervals for the KNN model across different partitions, including accuracy rates, confidence ranges, and AUC/Gini AUC scores, highlighting the model's overall efficacy and areas for potential improvement.

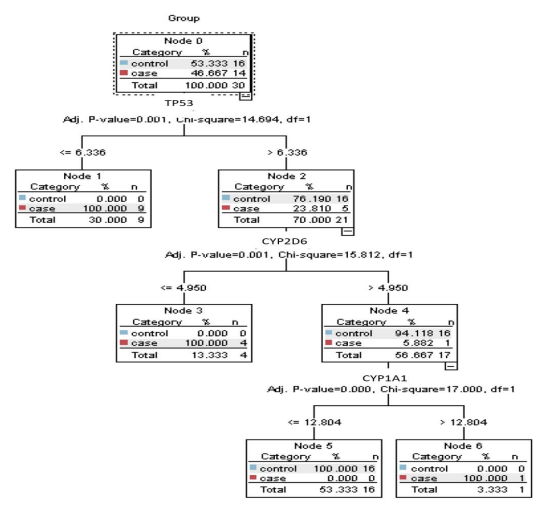
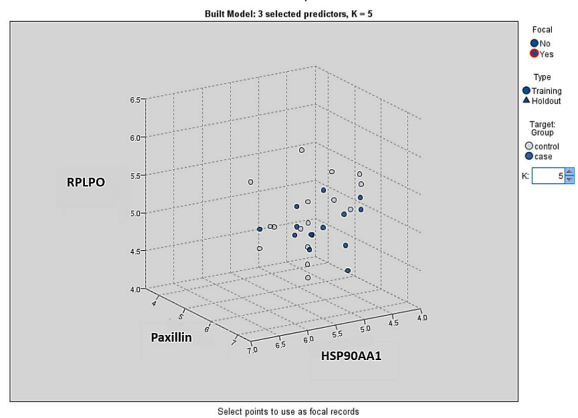


Figure 4. Decision tree generated by the CHAID algorithm

Predictor Space



This chart is a lower-dimensional projection of the predictor space, which contains a total of 414 predictors.

Figure 5. K-Nearest neighbors model predictor space (k=5) built on 3 selected predictors

Discussion

Despite advances in medical care and preventive measures in highincome countries, the overall incidence of stroke remains a substantial concern due to an aging population and associated risk factors (19). Given the limitations of current diagnostic methods, supplementary tools that provide timely and accurate information at the molecular level are increasingly necessary. Our findings advance previous research by demonstrating the effectiveness of RNA-seq data combined with machine learning models for ischemic stroke classification. Unlike traditional approaches that rely on neuroimaging, our method offers a molecular-level understanding of stroke pathology, identifying key genes (Such as TP53, CYP1A1, CYP2D6) and pathways (Including IL-17, NF-κB, and TNF signaling) that are typically not captured by imaging techniques. The identification of these pathways has important therapeutic implications, as they are central regulators of inflammatory and immune responses-critical components of ischemic stroke pathology. Targeting these pathways with anti-inflammatory agents or immunomodulators may provide new treatment opportunities that could improve clinical outcomes.

Recent advancements in genomics and bioinformatics have facilitated the discovery of novel biomarkers for stroke diagnosis and prognosis (20). These biomarkers, identified through high-throughput techniques such as RNA sequencing (RNA-seq), offer the potential for earlier detection and a deeper understanding of stroke pathophysiology (21). Parallel developments in machine learning (ML) and deep learning (DL) have introduced new strategies for enhancing diagnostic accuracy and treatment planning (22,23). These computational approaches can analyze large and complex datasets, identify subtle patterns, and produce predictions that may not be readily apparent through traditional statistical methods. ML algorithms, including Random Forests and KNN, have shown promise in medical applications such as stroke prediction and classification (24,25).

Machine learning has emerged as a transformative tool in medical research, particularly for analyzing complex data such as RNA-seq (26). ML algorithms can be used to classify stroke types, predict outcomes, and identify novel biomarkers by analyzing gene expression patterns and related features. The application of ML in stroke research provides several advantages. ML models can process large volumes of data and detect complex relationships that may be missed by traditional methodsan especially valuable capability given the high-dimensional nature of transcriptomic datasets (27). These models can be trained to predict stroke occurrence, severity, and outcomes using gene expression profiles, supporting early diagnosis and enabling personalized treatment strategies (28). ML methods also aid in dimensionality reduction, preserving essential biological information while improving model performance (29). Furthermore, ML models can integrate multi-omics data-such as genomics, transcriptomics, and proteomics-to provide a more comprehensive understanding of stroke pathology and identify therapeutic targets (30).

In this study, enrichment analyses (Figures 1 and 2) showed that the most prominent pathways were associated with inflammatory responses. We developed and validated several ML models using RNA-seq data to predict ischemic stroke, including Random Forest Classification, the CHAID Algorithm, and KNN. Each model offered unique insights into stroke prediction and classification, highlighting their specific strengths and limitations.

The Random Forest Classification model is a decision tree-based ensemble algorithm that builds multiple trees and aggregates their outputs to improve predictive accuracy (31). In our study, this model demonstrated exceptional performance, achieving 96.67% accuracy in the training set and 80% in the test set. The decision rules effectively classified stroke patients and controls based on gene expression thresholds. For example, certain rules indicated that high FOSB expression and low FAM83H-AS1 expression were associated with stroke. These findings are biologically plausible, as FOSB, a transcription factor, plays roles in cell proliferation, differentiation, inflammatory responses, and neuronal plasticity (32,33). FAM83H, while primarily known for its involvement in enamel formation (34), may influence inflammation or gene regulation, suggesting a possible role in stroke mechanisms.

The CHAID algorithm produced a decision tree with six nodes based on statistically significant splits (15). The model achieved perfect

accuracy in the training set and moderate-high performance in the testing set. The CHAID tree identified TP53, CYP1A1, and CYP2D6 as critical predictors. TP53 is a central regulator of apoptosis, inflammation, cell cycle progression, and DNA repair, making it highly relevant to ischemic injury and neuroprotection (35). CYP1A1, whose expression can be regulated by the aryl hydrocarbon receptor (AhR) and p53 pathways, may participate in oxidative metabolism and cellular responses to ischemia (36). CYP2D6, a major drug-metabolizing enzyme, may influence inflammatory processes or treatment responses in stroke patients (37,38). These genes therefore represent meaningful biological markers for classification.

The KNN model classifies samples based on similarity in the feature space (39). It demonstrated strong performance with high accuracy across training and testing partitions. The model identified RPLP0, Paxillin, and HSP90AA1 as significant predictors. RPLP0, a ribosomal protein involved in protein synthesis, may contribute to neuronal repair, though direct evidence in stroke is limited (40). Paxillin is involved in endothelial migration, inflammation, and vascular smooth muscle regulation-processes central to ischemic stroke pathology (41-43). HSP90AA1, a molecular chaperone, promotes neuroprotection, modulates inflammation, and may serve as a biomarker for stroke outcomes (42,44).

Overall, the ML models demonstrated strong potential for stroke prediction and classification. Each model offered distinct advantages. Random Forest provided clear and interpretable decision rules with high accuracy. The CHAID algorithm offered transparent decision paths based on statistical significance. KNN delivered strong predictive performance with high accuracy and confidence values. While KNN and Random Forest are computationally more demanding, all three models contribute unique perspectives on stroke prediction. The choice of model ultimately depends on research needs such as interpretability, accuracy, and computational resources. Combining multiple models could further improve predictive performance and offer more comprehensive assessments of stroke risk. Our findings align with previous studies that have used ML for stroke prediction, but our integration of RNA-seq data adds a novel molecular-level dimension.

Despite promising results, this study has limitations. The dataset consisted of only 40 samples (20 stroke patients and 20 controls), which may limit generalizability and statistical power. Additionally, model evaluation relied on a single train-test split without repeated randomization or cross-validation, which may introduce sampling bias and overestimate performance. Although our feature selection was focused on biologically relevant genes, incorporating additional omics data or clinical variables could improve predictive accuracy. Future research should validate these findings in larger, independent cohorts using rigorous validation methods such as k-fold cross-validation, and explore integrating molecular signatures into clinical workflows to support real-world diagnostic and therapeutic decision-making.

Conclusion

This study demonstrated the potential of machine learning techniques, including Random Forest Classification, the CHAID Algorithm, and K-Nearest Neighbors, in predicting and classifying ischemic stroke using RNA-seq data. Each model offered unique insights and strengths, highlighting the importance of ML in advancing stroke diagnosis and treatment. While the results are promising, further research and optimization are needed to enhance model performance and demonstrate their practical integration into clinical workflows.

The findings suggest that combining genomic data with advanced computational techniques can enable earlier detection and a deeper understanding of ischemic stroke mechanisms. Future studies should expand the dataset and incorporate additional omics layers to improve model robustness and generalizability. Ultimately, the application of ML in stroke research holds significant promise for improving diagnostic accuracy and personalizing treatment strategies, paving the way for more effective stroke management and better patient outcomes.

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Ethical statement

This study did not involve human participants, animals, or biological samples; therefore, ethical approval was not required.

Conflicts of interest

The authors declare no conflicts of interest relevant to this study.

Author contributions

Mina Rahmati: Data preprocessing, machine learning modeling, and drafting the manuscript. Masoud Arabfard: Supervision, statistical analysis, model evaluation, and manuscript revision. Both authors have read and approved the final manuscript.

Data availability statement

The raw RNA-seq gene expression data utilized in this study are publicly accessible in the Gene Expression Omnibus (GEO) repository under the accession number GSE22255. The analyzed datasets and associated computational findings are available upon reasonable request from the corresponding author, Masoud Arabfard.

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