ORIGINAL RESEARCH ARTICLE



A New Decision Tree to Solve the Puzzle of Alzheimer's Disease Pathogenesis Through Standard Diagnosis Scoring System

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Abstract Alzheimer's disease (AD) is a progressive, incurable and terminal neurodegenerative disorder of the brain and is associated with mutations in amyloid precursor protein, presenilin 1, presenilin 2 or apolipoprotein E, but its underlying mechanisms are still not fully understood. Healthcare sector is generating a large amount of information corresponding to diagnosis, disease identification and treatment of an individual. Mining knowledge and providing scientific decision-making for the diagnosis and treatment of disease from the clinical dataset are therefore increasingly becoming necessary. The current study deals with the construction of classifiers that can be human readable as well as robust in performance for gene dataset of AD using a decision tree. Models of classification for different AD genes were generated according to Mini-Mental State Examination scores and all other vital parameters to achieve the identification of the expression level of different proteins of disorder that may possibly determine the involvement of genes in various AD pathogenesis pathways. The effectiveness of decision tree in AD diagnosis is determined by information gain with confidence value (0.96), specificity (92 %), sensitivity (98 %) and accuracy (77 %). Besides this functional gene

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Department of Biotechnology and Bioinformatics, Jaypee University of Information Technology, Waknaghat, Solan, H.P. 173234, India classification using different parameters and enrichment analysis, our finding indicates that the measures of all the gene assess in single cohorts are sufficient to diagnose AD and will help in the prediction of important parameters for other relevant assessments.

Keywords Dementia · Alzheimer's disease · Mini-Mental State Examination · Classification · Decision tree · Clustering · Validation

1 Introduction

Alzheimer's disease (AD) is the most common form of escalating dementia in the elderly. It is a neurodegenerative disorder marked by the neuropathologic hallmark of intracellular neurofibrillary tangles (NFT) and extracellular amyloid plaques that accumulate in susceptible brain regions [1]. Difficulties remembering recent events are often early symptoms. Later symptoms include impaired communication, disorientation, confusion, poor judgment, behavioral changes and, ultimately, difficulty in speaking, swallowing and walking [2, 3].

AD is sixth dominating cause of death in the USA. Current statistics indicates that about 25–30 million people are afflicted from AD, and the number of cases will triple by 2050 due to increasing life expectancy [4]. According to 2015 population tally, an estimated 5.3 million American of all ages are suffered from this disease [5]. In addition to this, AD puts psychological and economical burden on caregivers and exhibits a major public health problem being among the most expensive disease at global level [6].

The amyloid and the tau hypothesis recognized amyloid precursor protein (APP) and tau proteins as inducers and key players of the disease [7]. Genes encoding these two



proteins have a varied influence on developing AD, ranging from the autosomal dominant inheritance in the familial forms (1–5 % of cases) to the polygenic background in late-onset (>65 years of age) and sporadic AD (=95 % of cases). In addition to the genetic component, the risk of developing AD is influenced by several other factors which include socio-demographic, lifestyle, environment and medical conditions [8]. Age and female sex represent varied risk of developing AD (Fig. 1).

The genetic component, however, seems to be of major concern, since according to twin studies, a major part of the risk of sporadic AD is genetically determined [9]. The APP, presenilin 1 (PSEN1) and presenilin 2 (PSEN2) genes are currently known to be involved in the familial forms of AD [10]. Identification and characterization of dominant mutations of these genes were auxiliary for the understanding of the biological mechanisms which lead to enhanced Aß accumulation and senile plaques formation [11]. In contrast to the familial AD, the causing factors of the Aß accumulations and other pathological mechanisms remain mostly unclear in the sporadic form. The complex genetic model of sporadic form suggests that several heterogeneous susceptibility sets of genes may converge on the pathological processes that underlie the disease [12]. However, so far only the apolipoprotein E (APOE) gene has been definitively associated with the risk of AD [13].

APOE is involved in lipid transport and metabolism. Furthermore, it plays a specific role in the central nervous system, including neuronal development, regeneration and certain neurodegenerative processes [14]. The polymorphism of the APOE gene determines three isoforms of APOE protein (e2, e3 and e4) with different conformation and lipid-binding properties [15]. A proportional relationship was found between the number of the inherited e4 alleles and the risk of developing AD and the age at onset. The APOE e4 isoform prefers very low-density lipoprotein, and it is less effective in cholesterol transport as compared

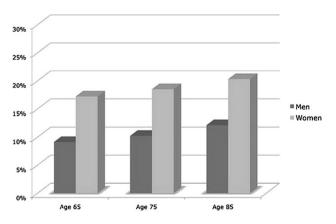


Fig. 1 Estimated lifetime risk of AD by age and sex ratio depicted in bar chart



to the other APOE isoforms. Membrane cholesterol modulates the cleavage of the APP protein, and in the presence of the e4 isoform, the balance is shifted to the production of Aß [16]. The amyloid cascade hypothesis has been the predominant model of molecular mechanisms underlying the pathogenesis of AD. According to this model, the genetic epidemiology of sporadic as well as familial AD remains a very active area of research, since a large part of the genetic etiology is still poorly understood and remains unresolved [17]. The aim of our work was to investigate direct entities and related attributes which are presumably involved in AD pathogenesis.

2 Materials and Methods

Classification of geneset data was carried out through RapidMiner Studio 6.2.0, Weka 3.6.9 and enrichment analysis for functional gene classification through David tool. Gene ontology (GO) study was done through webbased GEne SeT AnaLysis Toolkit (WebGestalt). All these analyses were performed on Window 7.0 platform running on a Lenovo PC with an Intel Core i5-2100 CPU processor and 4 GB of RAM.

2.1 Decision Tree

The decision tree represents tree-like structure to classify the data. Decision tree generates rules for classification [18]. A tree is represented by the set of nodes, leaves or branches. The root node is the attribute from which classification process starts, and the internal node corresponds to each of the questions about the particular attribute of the problem [19]. The branches coming out of the each node are labeled with possible values of attributes [20]. Each leaf node corresponds to a decision. The algorithm for generation of decision tree is partitioned into two parts: Top-down approach for induction of decision tree algorithm to choose features that partition the training data according to some evaluation function. Partitions are recursively split until some convergence criteria are reached. Secondly, the decision tree is pruned in order to avoid problem of over fitting [21] (Fig. 2).

The success of decision tree learning algorithm depends on evaluation criterion used to select the feature for splitting. Decision tree learning algorithm uses heuristic for estimating the best feature [22]. In our view, the measure yields a real positive number where the larger value indicates a set where there is more likelihood of all class values being present. Specifically, we use gain ratio (GR) as criteria for selection of features [23]. C4.5 algorithm implemented in RapidMiner tool treats missing value differently from normal values [24]. Association rules were generated

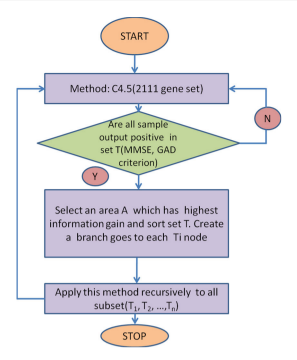


Fig. 2 Pipeline for decision tree building used for supervised classification of large gene dataset

which were determined by using Weka Tool, through various features.

2.2 Alzheimer's Gene Dataset

In this study, we have used Alzheimer's gene data available from various standard online resources such as ensemble gene, AlzGene, GenCard and NCBI. We selected 2111 raw genes that are known to relevant with Alzheimer's disease. Supplementary Table 1 contains list of genes used in our study. The dataset in our study consists of 14 attributes, which are describing different features of AD genes [25, 26]. Geneset was selected on the basis of the literature, and these genes were not only related to AD, in general, but also involved in other diseases. The most prioritizing attributes were selected based on the attribute selection techniques [27].

For selecting the attributes, the ranker technique was used. For the given dataset, the five different methods were applied. Chi-squared attribute Evaluation is the most widely used qualitative feature selection method. In order to reduce the effect of the bias resulting from the use of information gain, a variant known as gain ratio was utilized [28]. The gain ratio adjusts the information gain for each attribute to allow for the breadth and uniformity of the attribute values. Gain ratio is defined by the formula:

Gain ratio = information gain/intrinsic information.

The information gain attribute evaluation method evaluates the worth of an attribute by measuring the information gain with respect to the class.

Gain (Class, Attribute) = H (Class) - H (Class/Attribute),

where H is the information entropy.

2.3 Architecture and Model

The decision tree model is shown in Fig. 3. It begins with the collection of AD genes from different online resources, which is followed by preprocessing of the dataset including converting data into numeric values. Using geneset of 2111 gene, we use RapidMiner statistical tool and Weka Tool for data mining task. Further, we validated our model by using tenfold cross-validation application operator. Gene data include the descriptions such as gene name, association score, chromosomal position and MMSE score [29]. Enrichment analysis was also performed for this geneset to group genes on the basis of similarity using clustering algorithm. For more tightly associated gene in each group, stringency was maintained by keeping the kappa threshold 0.3 as anything below this threshold have great chance to be a noise. Finally, GO annotation was described through biological processes and molecular functions for the same.

3 Results and Discussion

Analysis was performed using classification of data by a J48 algorithm implemented in Weka, and C4.5 was used in RapidMiner. This analysis results in the correct classification of 950 out of 2111 genes. MMSE and Huge navigator were the most informative variable in this geneset obtained from information gain criterion. The classification algorithm provides MMSE score cutoff value for a different stage of the disease. We generated different rules through decision tree by selecting some specific parameter at each iteration [30].

The error rates and other features with respective classifier test for all decision trees are presented in Table 1.

After preprocessing the data which involve cleaning, integration and data reduction, the data mining was done. All analyses in this study involve tenfold cross-validation method to test the measures without pruning the tree. A single dataset of 2111 genes was used to build decision tree (DT). The dataset contains information about important attributes associated with genes. In order to improve the classifier, a supervised resample classifier was applied on data to assess the performance of the classifier. The number



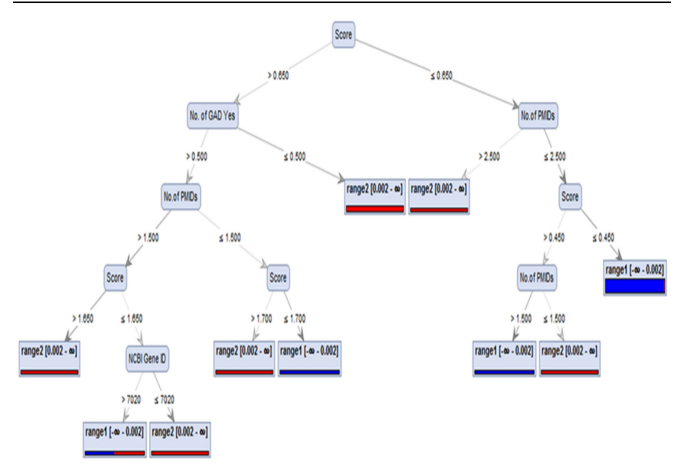


Fig. 3 Decision tree for the AD related genes. MMSE score is the root node at which classification of gene data has been done

Table 1 Statistical measure of performance of association rule (AR) based feature selection for sample group (AD genes dataset) using multiple setups

Data	Classifier	DT = 1	Features → error (%)			DT = 2	Feature → error (%))		
			F7	F11	F18	F32		F7	F11	F18	F32
2111 genes	C4.5	50	25.7	25.3	25.3	21.01	50	11.9	11.4	11.4	9.8
		100	23.9	23.2	23.2	20.4	100	9.8	9.5	9.5	7.4
		150	16.5	16.2	16.2	13.0	150	3.8	3.5	3.5	1.6

of tree was held constant at 100 while the number of features was kept varied at various points. Authenticity of built decision tree depends on accuracy which was measured 77 % with sensitivity 86 and specificity of the classifier was 81. On applying the priori algorithm, classifier generates ten best rules and was also verified (Tables 2, 3).

3.1 Decision Tree Induction

C4.5 algorithm implemented in the RapidMiner tool was used to make decision tree.

For making the decision tree, those feature were selected for classification whose values were not constant. Also we discretized the data on the basis of frequency. Different color node at the bottom of the tree reveals the class definition of leaf nodes.

3.2 Extracting Rule from RapidMiner

Based on the general process of data classification, we first identify a validation method to be used in modeling phase. The next step was to apply learning algorithm to training data and to generate rules. Decision trees were preferred since they are easy to understand. If the depth of tree increases, then IF–THEN rules can be extracted from decision tree.

To extract rule, every rule is drawn for each path from root node to the leaf node using some logical operators such as AND or IF. The summary table of decision tree (DT-1) is given below.



Table 2 Association between different classes based on a priori algorithm and best rule generated from classifier

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Minimum support 0.2					
Minimum confidence 0.9					
Number of cycle performed 1	6				
Best rules found					
a1 = false a5 = false 24	class = c0 24	conf:(1)			
$a5 = false \ a8 = false \ 24$	class = c0 24	conf:(1)			
$a5 = false \ a6 = false \ 23$	class = c0 23	conf:(1)			
a8 = false class = c1 22	a5 = true 22	conf:(1)			
a5 = false a7 = true 21	class = c0 21	conf:(1)			
$a5 = false \ a9 = false \ 21$	class = c0 21	conf:(1)			
$a3 = false \ a5 = false \ 20$	$class = c0 \ 20$	conf:(1)			
a6 = false class = c1 20	a5 = true 20	conf:(1)			
$a2 = false \ a5 = false \ 27$	$class = c0 \ 26$	conf:(0.96)			
$a4 = false \ a5 = false \ 23$	class = c0 22	conf:(0.96)			

3.3 Generation of Another Decision Tree Using CHAID Algorithm to Determine How Variable Best Combined Using Chi-Square Interaction Method

The performance of classifiers for both decision trees is calculated from the confusion matrix (Fig. 4; Table 4).

The first decision tree inputs only representative genes test samples accurately at 98.94 % using tenfold cross-validation strategy (Tables 5, 6).

Based on gene data, ten rules were generated for prediction of disease in DT-I and six rules in DT-II. On the basis of standard scoring system of MMSE score, performance of the model was improved (Tables 7, 8).

3.4 Data Analysis by Weka Tool

The AD gene dataset contains 499 relevant instances and 14 attributes. The data were analyzed using a WEKA software utilizing decision tree J4.8 classification algorithm and Bayesian network, and a Naïve Bayes algorithm. The classifiers were directly applied without any feature (gene)

Table 4 Summary of binary decision tree (DT) II

Learning algorithm	C4.5		
Attribute selection criterion	Gain ratio		
Input	2111 gene dataset		
Minimal gain	0.01		
Maximal depth	12		
Validation	Tenfold cross-validation		
Minimal split size	5		
Minimal leaf size	1		
Number of pre-pruning	3		

selection. The number of top-ranked genes selected using feature selection techniques and then classifiers technique was applied, on the data. The Relief Feature Attribute Evaluator is used in WEKA Explorer with a default parameter setting (Table 9).

Decision trees were generated by RapidMiner Studio and Weka Tool. Key attribute selection is a very critical step for the impactful decision tree. There are a bunch of methods available, but ranker search method is very powerful technique among these.

The performance of the classification model is measured by its predictive accuracy from the independent dataset. First, the classification model was built from a subset of the data called the training set, where the algorithm knows the values of both predictor attributes and classes for the data instances. After the model was built, its predictive accuracy was then measured in a separate subset of the data, called the test set, where the algorithm knows only the values of the predictor attributes (not classes) for data instances. So, this measure of predictive accuracy measures the generalization ability of the classification model and gave it better applicability.

3.5 Enrichment Analysis

3.5.1 GO Analysis

Enrichment analysis of the gene highlights the most relevant terms associated with GO with given gene list. Annotation includes GO which includes biological process,

Table 3 Summary table of decision tree (DT-I) with all essential parameters

Learning algorithm	C4.5				
Attribute selection criterion	Specifying the used method for selecting attributes, we choose gain ratio for this criterion				
Minimal size for split	4				
Minimal leaf size	1				
Minimal gain	0.1				
Maximal depth	20				
Confidence value	0.25				
Number of pre-pruning	3				



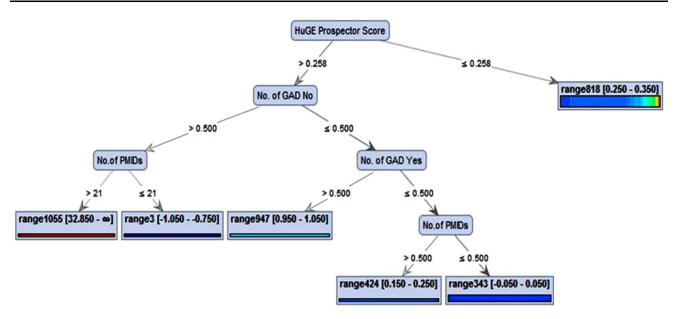


Fig. 4 Decision Tree of gene data based on Huge Prospector score as root node and number of GAD are important parameter for classification

Table 5 Confusion matrix for only representative 2111 gene for DT-I (without missing values)

	True range	True range	Class precision
Pred. range	1058	13	98.79 %
Pred. range	2	348	99.43
Class recall	99.81 %	96.40 %	

Table 6 Confusion matrix for only representative 2111 gene for DT-II (with missing values using CHAID method)

	True range	True range	Class precision
Prediction range	946	13	75.7 %
Pred. range	2	460	99.43 %
Class recall	75.7 %	83.5 %	

cellular component and molecular function [31]. The annotation coverage provides investigators with much more power to analyze their genes using different biological aspects in a single space. Annotation result for biological process and molecular function are displayed in the form of bar chart as shown in Figs. 5 and 6, respectively.

Members of independent groups fall into one of two mutually exclusive categories. Fisher's exact test was used to

Table 8 Class for building and using a 0-R classifier

Decision table summary	_		
Number of training instances	100		
Number of rules	31		
Start set	No attributes		
Search direction	Forward		
State search after five node expansions			
Total number of subsets evaluated	68		
Merit of best subset found	85		
Evaluation (for feature selection)	CV (leave one out)		
Feature set: 1, 6, 7, 9, 10, 11			
Correctly classified instances	77 %		
Incorrectly classified instances	23 %		
Kappa statistic	0.4912		
Mean absolute error	0.3316		
Root mean squared error	0.4002		
Total number of instances	100		

Predicts the mean (for a numeric class) or the mode (for a nominal class)

determine whether the proportions of those falling into each category differ by group [32]. To avoid over counting duplicated genes, the Fisher exact statistics is calculated based on

 Table 7 Decision

 table classifier output

	TP rate	FP rate	Precision	Recall	F-measure	ROC area	Class
	0.818	0.324	0.831	0.818	0.824	0.83	C0
	0.676	0.182	0.657	0.676	0.667	0.83	C1
Weighted avg.	0.77	0.275	0.772	0.77	0.771	0.83	



Table 9 Attribute-criterion-feature selection from the data on the basis of rank generated from the score

Attribute Evaluator	Supervised Filtered Attribute Evaluator			
Score	Attribute	Rank		
0.255714	a5	6		
0.038926	a3	4		
0.024319	a8	9		
0.009714	a2	3		
0.005152	a7	8		
0.003551	a9	10		
0.003551	a1	2		
0.002202	a6	7		
0.000531	a4	5		
0.000168	a0	1		

Feature selection is the technique of removing irrelevant features and to reduce dimensionality. The ranker search method is used to select attribute, and each selected attribute is ranked

corresponding ensemble gene IDs by which all redundancies in original IDs were removed. All results of chart report were generated after passing through the thresholds (by default, Max. Prob. ≤ 0.1 and Min. Count ≥ 2) [33].

The threshold of EASE score, a modified Fisher's exact *P* value, ranges from 0 to 1 and describes a statistically

Bar chart of Biological Process categories 1100 1000 900 800 Number of genes 700 600 500 400 300 200 100 metabolic process localization developmental process reproduction death cell proliferation growth unclassified oiological regulation response to stimulus nulticellular organismal process cell communication ellular component organization nulti-organism process

Fig. 5 Biological process under GO term; some biological phenomenon, commonly recognized series of events affecting state of an organism

significant number of genes in the list with respect to the number in the population of genes from which the list derives. Fisher's exact P value = 0 represents perfect enrichment. Usually, P value is equal to or smaller than 0.05 to be considered strongly enriched in the respective annotation categories.

3.5.2 Gene Functional Classification

For this purpose, hypergeometric statistical method was applied using Bonferroni method at significant level (0.05). From 2111 geneset, David tool found 958 valid IDs and 320 genes were irrelevant (supplementary file II). This classification of gene was based on important parameter as enrichment score which rank the biological significance of gene group based on overall EASE score of all enriched annotation terms. From 958 genes in DAVID tool, only 74 genes passed the filter and only this number of gene shows the common annotation term profile of functional group based on frequency [34]. Finally, 39 functional gene groups came as output on the basis of enrichment score range from 39.92 to 1.207 in decreasing order by applying certain parameter like kappa threshold of 0.3 and multiple linkage threshold of 0.50 as default setting (Supplementary file II).

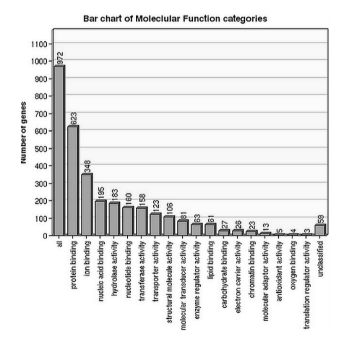


Fig. 6 Molecular function of involved gene under GO categories. The function carried out by a gene product; one product may carry out many functions; a set of functions together makes up a biological process



4 Conclusion

For human understanding, it is important to generate simple logic based classifier, i.e., in the form of a simple decision tree which describe the target concept. However, these simple decision trees may be of lower predictive value than other complex classifiers, but precision or accuracy generated from this classification model for gene dataset of 2111 genes is good (>80 %) while applying a C4.5 algorithm which follow branch and bound method for classification. The absolute error rate is very low. Specificity and sensitivity are also calculated to determine the suitability of designed model and are significant. Finally, from this study it is concluded that MMSE and relevance association score are important attributes for classification of genes and labeled them to a particular class. This type of testing and analysis has been used for selection of gene for expression arrays, automated protein data annotation, automatic cancer diagnosis, plant genotype discrimination, classifying gene expression profiles and computational model for mutational sites and then to extract best rules from designed models. Enrichment analysis alters us about the actual role of genes in term of GO, genes functional classification, pathway analysis, disease association, drug association and phenotype analysis. From gene functional classification analysis, we found APOE, PSEN1, GRN, ACE, BCHE, PRNP, IL1A are key genes that are strongly associated with AD whose association score ranges from 526.8 to 19.1 (Supplementary Table 1). Our proposed decision tree models and enrichment analysis of target genes will serve as a standard for computing biological phenomenon related to disease and exhibit their relevance toward AD conditions and its early diagnosis.

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