

Exploring Potential Biomarkers Underlying Pathogenesis of Alzheimer's Disease by Differential Co-expression Analysis

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Abstract

Background: Alzheimer's Disease (AD) is the most common form of dementia in the elderly. Due to the facts that biological causes of AD are complex in addition to increasing rates of AD worldwide, a deeper understanding of AD etiology is required for AD treatment and diagnosis.

Methods: To identify molecular pathological alterations in AD brains, GSE36980 series containing microarray data samples from temporal cortex, frontal cortex and hippocampus were downloaded from Gene Expression Omnibus (GEO) database and valid gene symbols were subjected to building a gene co-expression network by a bioinformatics tool known as differential regulation from differential co-expression (DCGL) software package. Then, a network-driven integrative analysis was performed to find significant genes and underlying biological terms.

Results: A total of 17088 unique genes were parsed into three independent differential co-expression networks. As a result, a small number of differentially co-regulated genes mostly in frontal and hippocampus lobes were detected as potential biomarkers related to AD brains. Ultimately differentially co-regulated genes were enriched in biological terms including response to lipid and fatty acid and pathways mainly signaling pathway such as G-protein signaling pathway and glutamate receptor groups II and III. By conducting co-expression analysis, our study identified multiple genes that may play an important role in the pathogenesis of AD.

Conclusion: The study aimed to provide a systematic understanding of the potential relationships among these genes and it is hoped that it could aid in AD biomarker discovery.

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Introduction

Aging causes an increasing susceptibility to cognitive performances due to a developing neurodegeneration leading to neurologic disorders, such as dementia. More than 20 million people worldwide suffer from dementia, and this number is expected to exceed 80 million by 2040 because of the rapid increase in the numbers of the elderly¹. Alzheimer's Disease (AD) is an irreversible progressive neurodegenerative disease affecting the central nervous system. Despite the increasing rate of AD incidence, no therapeutic strategy has been developed yet². Pathophysiologically, AD-related brain severe shrinkage caused neural and synaptic degenerations³. The mentioned degenerative events can be detected in post mortem examination of patients suffering from severe memory loss^{4,5}. It is thought that the loss of memory is because of aggregating beta amyloid (A β) and Neurofibrillary Tan-

gles (NFTs) of hyper-phosphorylated tau protein^{1,6}. Additionally, inflammation characterized by activated microglia⁷ and oxidative stress, which result from an imbalance of Reactive Oxygen Species (ROS) and antioxidants^{8,9} were shown to be associated with AD. Epigenetic changes happening in pre-frontal region by aging were shown to be related with AD functioning at cognitive level¹.

Rewiring of the biological networks to detect co-regulated and co-expressed units will help to facilitate looking into network's components and depicting the relationships between interconnected genes. Gene co-expression networks enable us to highlight molecular mechanisms underlying diseases¹⁰ and are considered as one way to investigate the etiology of AD efficiently. A large number of co-expression network methods have been proposed in the literature^{11,12}. Differential

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Co-expression Analysis (DCEA) offers a powerful approach for exploring phenotypic changes¹³. Not only is AD etiology incompletely understood but also differences at transcriptome level and the genes potentially related to each distinct regions of brain are not recognized causing AD to be remained somewhat unclear. In the present study, a high-throughput genomic screening approach was applied using DCGL software and comparative microarray analyses. It was hypothesized that the distinct transcriptional changes in different regions of brain lead to AD-associated brain damages. Therefore, the transcriptional profiles from the gray matter of frontal and temporal cortices were compared with hippocampi derived postmortem brains to dissect AD pathogenesis in these areas. The rationale behind the used network approach is to prioritize AD-causative genes that are apart from differential alterations in their expression and are differentially regulated by Transcription Factors (TFs) between contrasting samples. For this, Differential Regulation Analysis (DRA) has been conducted on three separated regions of AD brains as contrasting samples.

Materials and Methods

Data acquisition and pre-processing

The CEL files for GSE36980 series were downloaded from the GEO (<http://www.ncbi.nlm.nih.gov/geo/>) database and normalized with RMA method by using the Linear Models for Microarray Data (limma) R package. The main reason for selecting and exploiting this dataset is that GSE36980 series cover interspecies transcriptome analysis of various regions in gray matter in postmortem brains suiting the goal of dissecting pathological alterations in AD in several brain areas. Moreover, a number of researches have previously used these series and therefore would be able to compare the findings. After removing ambiguous probes, the extracted probe IDs were transformed into gene symbols. This data consists of a total of 79 samples (Table 1) based on the platform of GPL6244 and correspond to the frontal and temporal cortices and hippocampus.

Network construction

The DCGL R package was used to conduct DCEA^{13,14}. This software firstly calculates Differential Co-expression profile (DCp) and Differential Co-expression enrichment (DCE) to extract significant co-expression changes between a pair of genes in control and treatment samples. Next, Differentially Co-expressed Genes (DCGs) and Differentially Co-expressed Links (DCLs) were summarized from DCp and DCE values.

Next, DCGs and DCLs were extracted from DCp and DCE values previously calculated by DCp and DCE functions. DCp filters co-expression values of a pair of genes were assessed in control and treatment conditions. *X* and *Y* were defined as a subset of the gene pairs, where *n* is co-expression neighbors for a gene;

$$X = (x_{i1}, x_{i2}, \dots, x_{in})$$

$$Y = (y_{i1}, y_{i2}, \dots, y_{in})$$

The DC of a given gene is calculated with the following equation:

$$DC_n(i) = \sqrt{\frac{(x_{i1} - y_{i1})^2 + (x_{i2} - y_{i2})^2 + \dots + (x_{in} - y_{in})^2}{n}}$$

If the resulting DCGs and DCLs coincide with a TF, they will be referred to as a DRG and DRL, respectively. The DRGs and DRLs were scrutinized by DR-sort function in Differential Regulation Analysis (DRA) module. In fact, DRA module identifies potential TF as upstream regulators of DCGs and DCLs¹³. Finally, for illustrating the interactions between DRGs and their regulators, a network of DRGs and coincided TFs obtained by DRA was built for each of the datasets. By utilizing the Network Analyzer¹⁵ nodes were set within networks with higher connections to darker color and bigger size.

Gene ontology, pathway analysis and visualization

To find the significantly over-represented biological GO terms and functions of gene products within a co-expression network of DRGs and DRLs, functional classification was performed using BINGO Cytoscape plugin¹⁶ running hypergeometric test and Benjamini & Hochberg FDR correction at significant level 0.01. Finally, the clusters were visualized by Enrichment map Cytoscape plugin with Jaccard's coefficient 0.001. DRGs were further functionally classified by PANTHER database (<http://pantherdb.org/>) to underlying pathways (Figure 1).

Results

Co-expression analysis

The expression values of GSE36980 datasets were analyzed by utilizing DCGL v2.0 R package with default parameters. A total of 17088 unique genes were subjected to expression based filter and variance based filter, two functions embedded in DCGL to filter out genes that expressed extreme invariability across control and AD samples yielding 8544 and 2918 genes, respectively (Supplementary file 1). Afterward, using 2918 unique genes, co-expression analysis was performed on temporal cortex, frontal cortex, and hippocampus datasets separately. Expression based filter removes genes whose mean expression between experiments is lower than the median of this value for all genes and variance based filter removes genes that are not significantly variable than the median gene¹³. In order to prioritize seed genes which are potentially

Table 1. Sample characteristics

Biological samples	Control	AD patients
Temporal cortex	19	10
Frontal cortex	18	15
Hippocampus	10	7

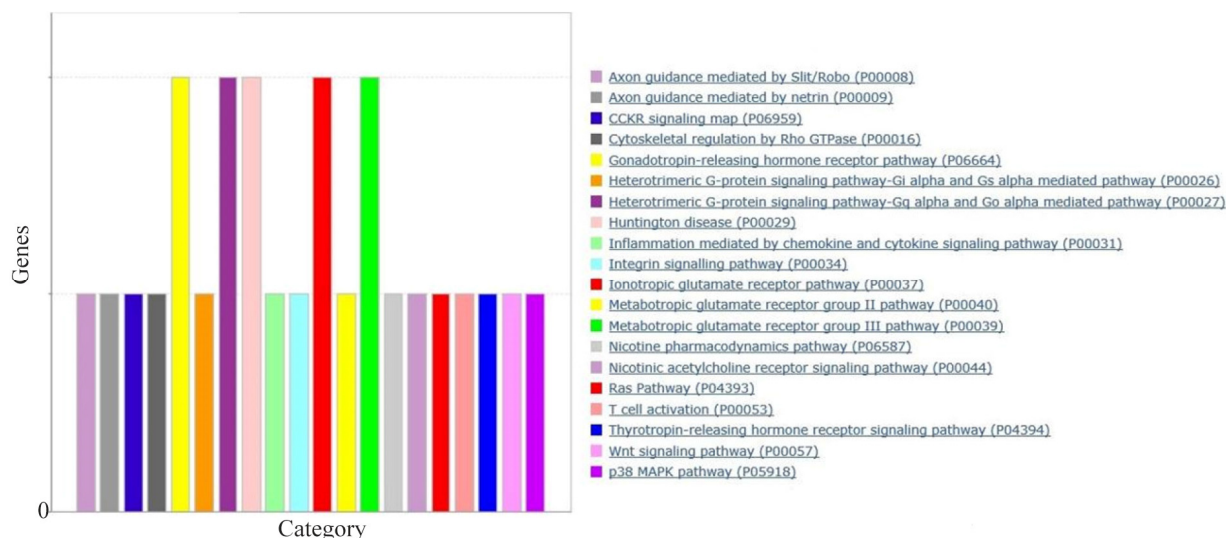


Figure 1. Bar chart of pathways potentially DRGs extracted from temporal cortex, frontal cortex and hippocampus expression data. PANTHER server with default parameters for pathway analysis was used for pathway analysis. The length of each bar shows how many genes have been assigned to a given pathway.

related to AD pathogenesis, common and significant DRGs were selected using Targets' Enrichment Density (TED) analysis and Targets' DCL Density (TDD) analysis. TED and TDD identify differential co-expression genes and link in a particular TF's targets, respectively¹³. To this end, targets of significant TFs were extracted from 19,9950 TF-to-target interaction pairs as a library in DCGL v2.0 software¹³. These pairs were further filtered out based on DRLs. In sum, 7, 19 and 13 genes were identified in temporal cortex, frontal cortex and hippocampus, respectively (Table 2). Significant TFs derived by TED and TDD analysis were used to infer co-expression network of DRGs in each dataset independently (Figure 2, Supplementary figures). DRGs were classified in terms of response to lipid, response to fatty acid, regulation of transcription from RNA polymerase II promoters and regulation of nitrogen compound metabolic process (Figure 3). Moreover, in pathway analysis, signaling pathways such as glutamate and G-protein signaling pathways were noteworthy (Figure 1).

Temporal cortex

460 DCGs and 33656 DCLs were summarized using DC sum function to a final set of DCGs and DCLs (Supplementary file 2). There were 199 significant TFs in the results of TED analysis and 35 significant TFs in TDD analysis. 35 TFs that were significant in both of these two analysis results were chosen (Supplementary file 2). DRA analysis yielded 7 DRGs and 33 DRLs. DRGs were not only differentially co-expressed but also differentially co-regulated with 35 mentioned TFs. Then, a network of DRGs and DRLs was visualized using Cytoscape 3.4.0. Based on figure 2, *PAX5* transcription factor and genes including *ARID1A*, *CDC42* and *LPPR4* were highlighted as the most important

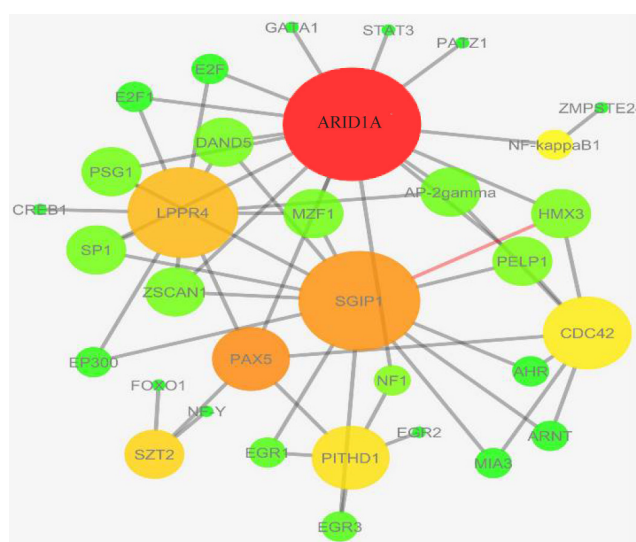


Figure 2. Differential co-expressed network of DRGs and DRLs captured by TED and TDD results in temporal cortex datasets. The bigger and darker nodes show the nodes with higher connectivity within the network.

units within the genes network with more interconnected links (Figure 2).

Frontal cortex

In frontal cortex datasets, 628 DCGs and 166256 DCLs were summarized to 20 DRGs and 164 DRLs (Supplementary file 3). There were 199 significant TFs in TED analysis result and 135 significant TFs in TDD analysis result from which 135 TFs were chosen that were significant in both TED and TDD results (Supplementary file 3). In the inferred network, *PAX5* and *IKZF1* as TFs and genes including *GRIK3*, *MAGI3*, *PRRX1* and *DCAF6* were found as highlighted nodes with more connectivity.

Biomarkers Underlying Pathogenesis of Alzheimer's Disease

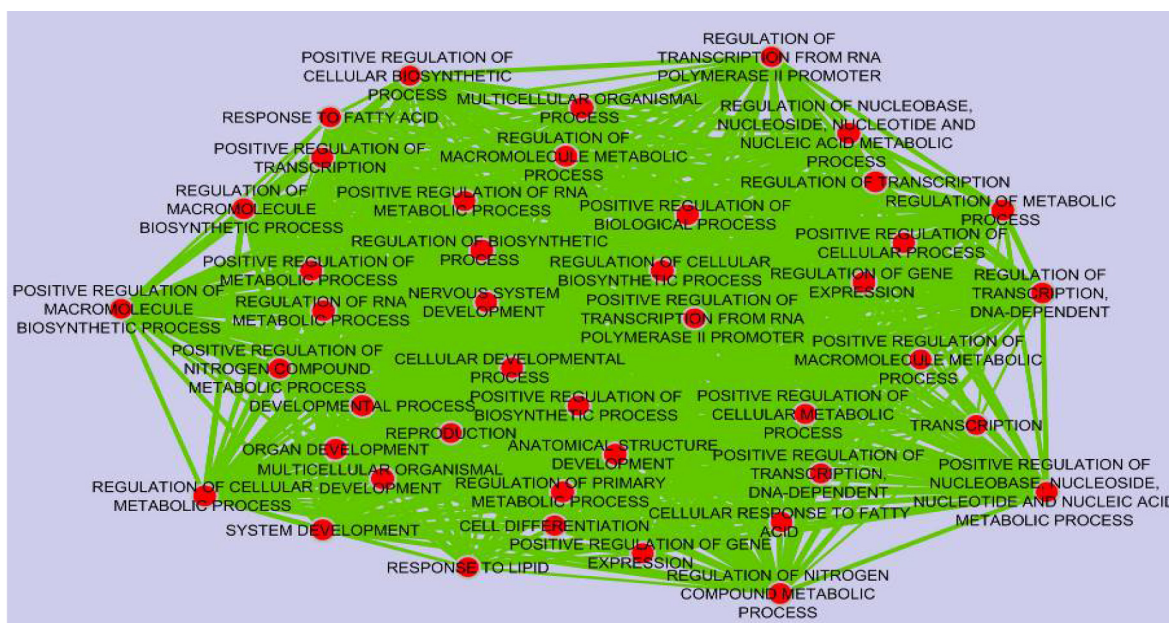


Figure 3. Functional classification of biological process in which Differential Regulated Genes (DRGs) were supposed to be involved. The GO terms were considered significant based on hypergeometric test with Benjamini & Hochberg FDR correction and significance level 0.01 by BINGO app. The results were illustrated using the Enrichment map Cytoscape 3.4.0 plugin. Ticker lines and bigger circles show more genes with higher significance level belonging to a given term.

Hippocampus

According to hippocampus datasets, 670 DCGs and 56264 DCLs were summarized to 16 DRGs and 43 DRLs (Supplementary file 4). There were 199 common and significant TFs in TED and TDD analysis which were used for inferring differential co-expression network with DRGs. There was more connectivity in hippocampus network than the other two networks. *PAX5*, *ARNT*, *GATA1*, *EGR3* and *IKZF1* TFs and genes including *KCNK1*, *CACHD1*, *FABP3* and *CHRNA2* showed highlighted roles as network nodes.

Discussion

Aging is believed to be one of the most important non-modifiable risk factors of cognitive diseases that lead unequivocally to a number of detrimental changes in the neural system, increasing neuromorbidity and mortality. AD, as a progressive neurodegenerative disorder with no effective treatment options, is typically characterized by the presence of amyloid-beta plaques and hyper phosphorylated paired helical filament tau protein-rich neurofibrillary tangles¹. The identification of co-expressed genes related to AD presumably provides insights into the underlying mechanisms; in other words, a combination of gene effects likely holds promise as a more effective approach for detecting disease associated genes⁴². In fact, examining co-expressed genes in spite of the individual genes could be more informative to explore genes that cause mental health disorders, such as AD^{43,44}. In this case, the correlation between two genes varies in distinct samples and thereby they are referred to as being differentially co-expressed.

This correlation may change independently from the expression levels of two genes, indicating that transcriptome analysis merely based on differential expression analysis could miss important clues of regulatory patterns⁴⁵. Co-expression analysis has been performed for deciphering molecular mechanisms underlying mental health disorders⁴⁶⁻⁴⁹. In the context of a well-established network analysis approach and given the most variable transcripts between control and AD brain samples, attempts were made by DCGL framework to explore putative pivotal genes that may be associated with AD. This work attempted to identify DRGs and links DRLs in AD by comparing expression datasets of temporal and frontal cortices and hippocampi. A comprehensive search in the literature showed that the obtained DRGs of AD brains mostly have direct or indirect links with AD or another neurologic disorder (Table 2). They are implicated in the gene ontology terms and shared biological pathways like response to lipid, fatty acid, nitrogen compound metabolic process and glutamate signaling pathways (Figures 1 and 2, supplementary file 5). Reportedly, considering GO terms as the response to lipid and fatty acid, brain lipid homeostasis plays an important role in AD⁵⁰. In this regard, differential regulation of delta 4-desaturase, sphingolipid 1 (*DEGSI*) and fatty acid binding protein 3 (*FABP3*) in hippocampus and lipid phosphate phosphatase-related protein type 4 (*LPPR4*) in temporal cortex datasets may fairly explain the relationship between brain damages happening in these regions and lipid metabolism. *DEGSI* encodes a member of the membrane fatty acid desaturase family which is shown

Table 2. List of differential regulated genes (DRGs) and corresponding p-value<0.05 of differential co-expression enrichment (DCe) in temporal cortex, frontal cortex and hippocampus datasets

	Gene name	DCe p-value	Description
Temporal cortex	<i>ARID1A</i>	0.00069	<i>ARID1A</i> was among down-regulated genes in AD model mice (17)
	<i>Cdc42</i>	0.01224	<i>Cdc42</i> activity was increased in hippocampus neurons treated with fibrillary β -amyloid (18)
	<i>LPPR4</i>	0.01395	<i>LPPR4</i> was up-regulated in incipient AD patients (19)
	<i>PITHD1</i>	0.01863	//
	<i>SGIP1</i>	0.02064	//
	<i>SZT2</i>	0.01936	//
	<i>ZMPSTE24</i>	0.01936	//
Frontal cortex	<i>CHD5</i>	5.19E-14	The depletion of <i>CHD5</i> was shown to be linked with AD associated gene sets (20)
	<i>EFHD2</i>	3.00E-07	<i>EFhd2</i> has been found to be associated with aggregated tau in the brain in AD and in a mouse model of frontotemporal dementia (21,22)
	<i>Prxs</i>	7.82E-07	Peroxiredoxins (Prxs) may be associated with AD by reducing ROS elicited by amyloid β (A β) accumulation that could be a causative factor in the pathogenesis of AD (23)
	<i>MAGIE3</i>	1.68E-06	//
	<i>EXTL1</i>	4.14E-06	//
	<i>HPCAL4</i>	2.09E-05	<i>HPCAL4</i> could be used as a prognostic marker for cognitive decline in AD (24)
	<i>LPHN2</i>	2.59E-05	<i>LPHN2</i> is likely to be participated in AD as an altered protein in Lipid Raft (25)
	<i>NIPAL3</i>	8.35E-05	<i>NIPAL3</i> was shown as a biomarker in Late-Onset Major Depressive Disorder (26)
	<i>CACNA1E</i>	0.00017	<i>CACNA1E</i> was down-regulated in cerebral Cockayne syndrome (27)
	<i>IFI16</i>	0.00033	<i>IFI16</i> was participated in delaying onset of AD (28)
	<i>HHLA3</i>	0.00122	//
	<i>KCNK1</i>	0.00202	<i>KCNK1</i> exhibited alternative splicing in patients with mesial temporal lobe epilepsy (29)
	<i>rnp3</i>	0.00384	//
	<i>DCAF6</i>	0.00542	//
	Hippocampus	<i>IPO13</i>	0.00581
<i>RPL11</i>		0.00585	<i>RPL11</i> revealed significant altered expression profiles in the neuron model of AD treated with rhTFAM (31)
<i>S100A1</i>		0.00716	<i>S100A1</i> modulates inflammation in AD (32)
<i>CNTN2</i>		0.02606	<i>CNTN2</i> associated with AD via <i>BACE1</i> activity (33)
<i>GRIK3</i>		0.03774	<i>GRIK3</i> was highly expressed in major depression (34)
<i>KCNK1</i>		1.55E-09	//
<i>CHRNA2</i>		2.37E-09	<i>CHRNA2</i> was found to interfere with the immune system in neurologic disorders (35)
<i>HAPLN2</i>		3.43E-05	<i>Hapln2</i> has been recently shown to be accumulated in the neurofibrillary tangle of Alzheimer's brain (36)
<i>Slc2a1</i>		0.00207	<i>Slc2a1</i> down-regulation exacerbated AD (37)
<i>FABP3</i>		0.00298	serum levels of brain-type <i>FABP</i> are elevated in a significant proportion of patients with various neurodegenerative diseases including AD (38)
<i>DEGS1</i>		0.00327	<i>DEGS1</i> is likely to be involved in AD as an altered protein in Lipid Raft (25)
<i>NKAIN1</i>		0.00435	//
<i>S100A1</i>		0.00434	<i>S100A1</i> modulates inflammation in AD (32)
<i>CNTN2</i>		0.00511	<i>CNTN2</i> associated with AD via b-Secretase (<i>BACE1</i>) activity (33)
<i>SFPQ</i>		0.00816	<i>SFPQ</i> was shown as a transcription factor with an altered nucleo-cytoplasmic distribution under neurodegenerative conditions (39)
<i>GPSM2</i>	0.01073	//	
<i>GSTM1</i>	0.01591	<i>GSTM1</i> null genotype was found as risk factor for late-onset Alzheimer's disease in Italian patients (40)	
<i>CACHD1</i>	0.02830	<i>CACHD1</i> is a substrate of <i>BACE1</i> responsible for generating the amyloid-b protein (41)	

// showing DRGs with ambiguous role in neurologic disorders.

to interfere in AD via lipid rafts²⁵. FABP proteins are thought to participate in the uptake, intracellular metabolism and/or transport of long-chain fatty acids. Concordantly, serum levels of brain-type *FABP* are elevated in a significant proportion of patients with various neurodegenerative diseases including AD³⁸. *LPPR4* acts as phospholipid dephosphorylate involving axonogenesis. The control of ion flow across the lipid

membrane is essential for many cellular functions, including neuronal excitability and dysfunction of conveying ions through lipid bilayers involved in multiple neurologic diseases⁵¹. As illustrated in figure 1, the DRGs are more implicated in signaling pathways; but the DRGs from frontal cortex were more enriched in ionotropic glutamate receptor pathway and metabotropic glutamate receptor group II and III pathways.

The dysregulation of glutamatergic signaling has been shown to be associated with AD. Glutamate acts *via* ionotropic glutamate receptors (*iGluR*) and metabotropic glutamate receptors (*mGluR*), both of which have been implicated in AD⁵². Differential regulation of glutamate receptor ionotropic, kainate 3 (*GRIK3*) and voltage-dependent R-type calcium channel subunit alpha-1E (*CACNA1E*) in frontal cortex datasets may be biologically relevant with the mentioned pathways in AD brain areas. Concordantly, a significant change in the expression of the *GRIK3* gene was detected in a patient diagnosed with severe developmental delay⁵³. Many different kinds of signaling pathways are changed in AD, indeed the relevance of the biological pathways shown in figure 1 such as cytoskeletal regulation by Rho GTPase suggests mediating of these signaling pathways in the different lobes of brain, in this case in temporal cortex with differential regulation of *CDC42*. *CDC42* has been linked to neuronal diseases like Alzheimer and Parkinson's disease through its role in cytoskeletal organization⁵⁴. Among the DGRs, *CNTN2*, *KCNKI* and *S100A1* were found common in frontal cortex and hippocampus datasets. *S100A1* encodes for calmodulin signaling molecules. Increased levels of calmodulin have been reported in the hippocampus of AD model mice⁵⁵. These changes seemingly show an aberrant involvement of calmodulin in the impairment of cell cycle control in AD. As for the potassium channel subfamily K member 1- *KCNKI*, recent genetic studies suggest a central role for neuroinflammation. *KCNKI* is a voltage-gated potassium channel upregulated by activated microglia and a mediator in amyloid-mediated microglial priming, additionally reactive oxygen species production that was shown to be related with autoimmunity⁵⁶. *CNTN2* has been shown to undergo nuclear translocation and altered transcription³³.

These findings probably show that hippocampus and frontal cortices might deeply play a role in AD by mediating with conveying ions. Their obtained DRGs participated in vital processes like signaling, ion transportation and homeostasis. However, these processes mostly signal pathways somehow shared with temporal cortex implying the role of signal molecules within and between brain areas in neurologic dysfunctions. Concordantly, a comprehensive study has been already carried out on GSE36980 series to examine the alteration in the expression of diabetes-related genes in AD brains where they illustrated that hippocampi of AD brains have the most significant alteration in gene expression profile⁵⁷.

With a glance at table 2 and the terms including amyloids, inflammation, ROS and immune system, one could infer a cascade of events in which the DRGs interfere. Beta-amyloid deposition following the activation of microglia will initiate an inflammatory response leading to the release of potentially neurotoxic substances and ROS that targets neural damage⁵⁸. Afterward, along with immune response, nitrogen com-

pounds will mediate to reverse the consequences of oxidative stress in damaged regions^{8,9}. In sum, it was shown that DRGs covered a wide range of known functions and processes implicated in main AD signaling pathways. In a study by Satoh *et al*⁵⁹, GSE36980 series used in the present study were utilized to identify biomarker genes relevant to the molecular pathogenesis of AD. They analyzed a RNA-Seq dataset composed of the transcriptome of postmortem AD brains derived from two independent cohorts and they identified the core set of 522 genes deregulated in AD brains shared between both, compared with normal control subjects. Notably, in agreement with our study, *LPPR4* was bolded in AD brains in both microarray and RNA-seq datasets. By consistent downregulation of *NeuroD6* in AD brains, the results indicated that downregulation of *NeuroD6* serves as a possible biomarker for AD brains. Previous studies identified *LPPR4* as direct target genes for *NeuroD6* by binding assay to E-boxes located in target gene promoters⁶⁰. GSE36980 series were also employed by Fowler *et al*⁶¹ used to investigate potential underlying biology in AD and in concordance with the results of the present study, they noticed the overrepresentation of glutamate in their data mining. They first identified genes consistently associated with AD in each of the four separate expression studies, and confirmed the result using a fifth study. They next developed algorithms to search hundreds of thousands of GEO data sets, identifying a link between an AD-associated gene (*NEUROD6*) and gender. Additionally, they identified several genes related to glutamate (including *CACNG3*, a regulator of AMPA-sensitive glutamate receptors; *SLC17A7*, a mitochondrial oxoglutarate carrier; and *GOT2*, mitochondrial glutamic-oxaloacetic transaminase. In our study, differential regulation of glutamate receptor ionotropic, kainate 3 (*GRIK3*) and voltage-dependent R-type calcium channel subunit alpha-1E (*CACNA1E*) in frontal cortex datasets could be therefore biologically relevant with the mentioned pathways in AD brain areas. Moreover, in our study, differential regulation of *Slc2a1* in hippocampus data seemingly implies the role of impairments in glutamatergic transmission mostly in hippocampus of AD brains. The role of glutamate transporters such as *SLC1A6* was also highlighted in a study by Satoh *et al*⁵⁹.

Conclusion

The purpose of the study was to explore the molecular mechanism in the development of AD, and a comparison of AD in three regions of the brain was done. Therefore, in the frame of network reconstruction and data mining approaches, a small number of possible genes and TFs were identified that their interplay could lead to neural dysfunctions toward AD. However, one should be cautious regarding small sample size while by utilizing more adequate samples, the results would be more reliable evidences.

An expected outcome of such a work would possibly shed light on the bridges between AD-associated brain damage in transcriptome level and presenting crucial evidence in clinical diagnosis and treatment.

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Conflict of Interest

The authors declared that they have no competing interests.

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