



FULL LENGTH ARTICLE

Gene interactions analysis of brain spatial transcriptome for Alzheimer's disease

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Abstract Recent studies have explored the spatial transcriptomics patterns of Alzheimer's disease (AD) brain by spatial sequencing in mouse models, enabling the identification of unique genome-wide transcriptomic features associated with different spatial regions and pathological status. However, the dynamics of gene interactions that occur during amyloid- β accumulation remain largely unknown. In this study, we performed analyses on ligand-receptor communication, transcription factor regulatory network, and spot-specific network to reveal the dependence and the dynamics of gene associations/interactions on spatial regions and pathological status with mouse and human brains. We first used a spatial transcriptomics dataset of the *App^{NL-G-F}* knock-in AD and wild-type mouse model. We revealed 17 ligand-receptor pairs with opposite tendencies throughout the amyloid- β accumulation process and showed the specific ligand-receptor interactions across the hippocampus layers at different extents of pathological changes. We then identified nerve function related transcription factors in the

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hippocampus and entorhinal cortex, as well as genes with different transcriptomic association degrees in AD versus wild-type mice. Finally, another independent spatial transcriptomics dataset from different AD mouse models and human single-nuclei RNA-seq data/AlzData database were used for validation. This is the first study to identify various gene associations throughout amyloid- β accumulation based on spatial transcriptomics, establishing the foundations to reveal advanced and in-depth AD etiology from a novel perspective based on the comprehensive analyses of gene interactions that are spatio-temporal dependent.

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Introduction

The relationship of amyloid plaques to the neurodegenerative process and related gene expression changes is a central question in Alzheimer's Disease (AD) research. Spatial transcriptomics (ST) is an attractive approach that provides unbiased transcriptome profiling of sequenced molecules with spatial localization and focused image signals like amyloid plaques.^{1,2} Previous studies used mouse brains from *App*^{NL-G-F} (AD model) and C57BL/6 (wild-type/WT) mice at 3, 6, 12, and 18 months of age to explore ST with the corresponding quantitative amyloid plaque index information, and demonstrated alterations in the plaque-induced and oligodendrocyte gene networks in different phases of AD. However, the dynamic changes of gene interactions between ligand-receptor (L-R) pairs, mediated by transcription factor (TF) regulation, and various degrees of gene associations/interactions at the transcriptome level throughout the disease process in different brain regions remain largely unexplored.

Defining a more detailed landscape of how different types of gene interactions are altered in the neuronal structures of the brain may contribute to a better understanding of AD etiology.^{2–4} To accomplish this goal, multiple approaches such as CellPhoneDB⁵ and single-cell regulatory network inference and clustering (SCENIC)⁶ have been used to resolve spatio-temporal differences in gene interactions in the same ST dataset. In addition, ST allows gene expression profile selection to be anchored to some particular brain regions of interest, which in this study were the hippocampus (HP) and entorhinal cortex (ENT). The HP and ENT play central roles in AD pathology. Hippocampal deficits in amyloid- β (A β)-related rodent models of AD have revealed synaptic, behavioral, and circuit-level defects.^{7,8} ENT is one of the earliest affected brain regions.⁹ A β proteins are seen primarily in the ENT in mild AD and "spread" to the HP and other cortical areas as the disease progresses.¹⁰ Selective overexpression of mutant amyloid precursor protein (APP) in ENT has been reported to cause an aberrant excitatory cortico-hippocampal network activity leading to behavioral abnormalities.^{10,11}

Here we used the ST dataset of *App*^{NL-G-F} and C57BL/6 mice with annotations for brain region,¹² AD phenotype, and A β index to explore the genome-wide transcriptomic features associated with different spatial regions and pathological status. In general, our work revealed 17 L-R pairs with opposite tendencies throughout the A β accumulation process. We also showed the specific L-R interactions across the HP layers at different extents of

pathological changes which may be related to their unique functions cross these region pairs. We searched the convergent functional genomic (CFG) scores of these unique L-R pairs in the AlzData database¹³ to validate their AD-related functions. Next, we identified nerve function related TFs in the HP and ENT in order to explore their changes with disease progression. We then calculated genes with different degrees (interactive connection with other genes) based on the network degree matrix (NDM) from spot-specific network (SSN) analysis. The purpose was to reveal unique gene connections in different brain regions and age groups of different phenotypes (WT/AD). These results identified novel and known L-R pairs/genes that may contribute to disease pathology and revealed the strong and coordinated cellular responses in the amyloid plaque niche based on a comprehensive analysis of gene interactions that are spatio-temporal dependent.

Materials and methods

Data sources and workflow

The whole brain ST dataset used in the present study was obtained from the Gene Expression Omnibus (GEO) database (<https://www.ncbi.nlm.nih.gov/gds>) with accession number GSE152506. This dataset¹² integrated 10,327 transcriptomics profiles (10,327 spots/tissue domains) from *App*^{NL-G-F} and C57BL/6 male mice at 3, 6, 12, and 18 months of age as one gene expression matrix annotated with Brodmann area, pathological status (A β index), and age information. Twelve mouse samples were used in this dataset: mouse number (n) = 2 per phenotype (WT or AD) for 3 and 18 months of age; n = 1 per phenotype (WT or AD) for 6 and 12 months of age; n = 12 for total. Two ST slides were included per brain sample from each of the 3- and 18-month mice. Only one ST slide was included per brain sample from each of the 6- and 12-month mice. The total ST slide number was 20, including 4 in WT_03 (3 months), 1 in WT_06 (6 months), 1 in WT_12 (12 months), 4 in WT_18 (18 months), 4 in AD_03, 1 in AD_06, 1 in AD_12, and 4 in AD_18. Spatial transcriptomics experiments were performed following the Library Preparation Manual (Spatial Transcriptomics, Stockholm, Sweden).¹⁴ The diameter of a tissue domain is 100 μ m. Data integration and normalization (by the EdgeR "cpm" function) for the multiple slides for the same age and phenotype groups were done in the original paper¹² as well as the log-cpm matrix downloaded from GEO.

The HP ST dataset used in the present study was available at Mendeley <https://doi.org/10.17632/6s959w2zyr.1y>. This dataset came from a study² which included two types of AD models. The triple transgenic AD model (3xAD) expresses three human gene variants: APP K670N/M671L, MAPT P301L, and PS1 M146V. 3xPB mouse is a modified version of the 3xAD strain formed by adding a deficiency in DNA polymerase beta (Polb). The 3xPB AD model displays more severe deficits in memory, learning, and long-term potentiation as well as higher levels of DNA damage and cell death. All mice are male (11–13 months of age) on a C57BL/6J background. For HP ST, three mice per model and two sections per mouse were included.

The ENTI single-nuclei RNA sequencing (snRNA-seq) dataset used in the present study was obtained from the GEO database with accession number GSE138852. This dataset came from a study¹⁵ in which post-mortem AD and non-diseased age-matched human ENTI tissues were collected from the Victorian Brain Bank. Disease or control group allocation was based on overall amyloid and tau pathology. In total, this dataset included six AD and six control gene expression matrices (6 AD patients and 6 control individuals) from snRNA-seq with annotations for cell type and phenotype.

The functions of genes in significant L-R pairs or TF regulatory networks were validated in an AD database, AlzData (<http://www.alzdata.org/>).¹³ AlzData provides various types of information for target genes, including the expression of the target gene regulated by AD genetic variants; significant physical interactions of proteins of the target genes with AD core genes such as APP, PSEN1, PSEN2, APOE, or MAPT¹³; and differentially expressed in AD mouse models before AD pathology emergence and the correlation of target gene expression with AD pathology in amyloid beta or tau line AD mouse models. One CFG score is added if any of the above evidence is significant. The CFG score ranged from 0 to 5.

In brief, we used the whole brain ST dataset (GSE152506) to perform L-R interaction analysis within spots and across brain regions (HP layers). Next, TF network and SSN analysis were performed in HP and ENTI areas. Then AlzData database was used to validate the target genes' AD-related functions by the CFG score. We also used the independent HP ST dataset to validate the expression of unique L-R interactions that are only differentially expressed in one HP region pair in the whole brain ST dataset. The ENTI snRNA-seq dataset was used to validate the different expression of important TFs in human AD patients compared with the control (Fig. 1A).

Analysis of L-R communication

We used CellPhoneDB,⁵ a public knowledge base of ligands-receptors that considers interacting partners as binary interactions annotated by IUPHAR (<http://www.guidetopharmacology.org/>) and cytokines, hormones, and growth factors interacting with receptors annotated by the iMEX consortium (<https://www.imexconsortium.org/>) to systematically analyze the intra-spots/cross brain regions in their communication network, and calculate the average log gene expression level and communication significance of each L-R pair. To perform statistical inference of L-R

specificity, a null distribution for each L-R pair mean was generated by random permutation of spot labels (1000 times by default). By calculating the proportion of the means that are "as or more extreme" than the actual mean, CellPhoneDB obtained a *P*-value for the likelihood of spot-type specificity of a given L-R pair. Predicted interaction pairs with *P*-values <0.05 were considered as significant as defined in the original analyses.⁵

The intra-spot molecular interactions analysis only used CellPhoneDB to get the expression value of each L-R pair within each spot. Specifically, CellPhoneDB only provided the L-R database and calculated the average log gene expression level of ligand and receptor genes in the same spot in this part.

In the cross-region L-R analysis, CellPhoneDB compared the average log gene expression level of each L-R pair among different region pairs. *P*-values <0.05 were considered statistically significant (Fig. S1A). The output results (Fig. 4A, B) were the total significant L-R pair numbers (Fig. S1B) in each pair of regions.

Our L-R communication analysis needed more detailed subgroup classification (regarding certain brain regions in the same age, A β index, and phenotype group) information than TF regulatory network and SSN analyses as input. However, the expression data missing rate in some brain regions at certain A β accumulation levels was higher in 6- and 12-month-age groups than the two other age groups. This is because the total numbers of spots in these groups were limited in the whole brain ST dataset compared with the 3- and 18-month-age groups. So, the 6- and 12-month-age groups were not involved in our L-R communication analysis.

Soft clustering analysis

Soft clustering methods such as Mfuzz can assign genes to several clusters using the fuzzy c-means algorithm with time/process-course data on the gene expression and may be more accurate and robust than hard clustering.¹⁶ To analyze the expression trend of intra-spot L-R pairs with A β accumulation, the Mfuzz package in R was used to perform soft clustering analysis for visualizing the A β level-dependent expression patterns of L-R pairs and assign L-R pairs to several cluster modules with different expression tendencies through A β accumulation. Parameter *m* was calculated by the "mestimate" function.

Gene enrichment analysis

To identify the significantly enriched pathways of significant L-R gene pairs, we used the clusterProfiler R package to perform Gene Ontology (GO) enrichment analysis. Only terms showing adjusted *P*-values less than 0.05 (adjusted for multiple testing by using the BH method) were considered as significantly enriched.

SCENIC analysis

First, putative TF motifs were identified by RcisTarget (hg19-tss-centered-10kb-7species.mc9nr.feather, <https://resources.aertslab.org/cistarget>) software in Bioconductor. The SCENIC algorithm was then used to

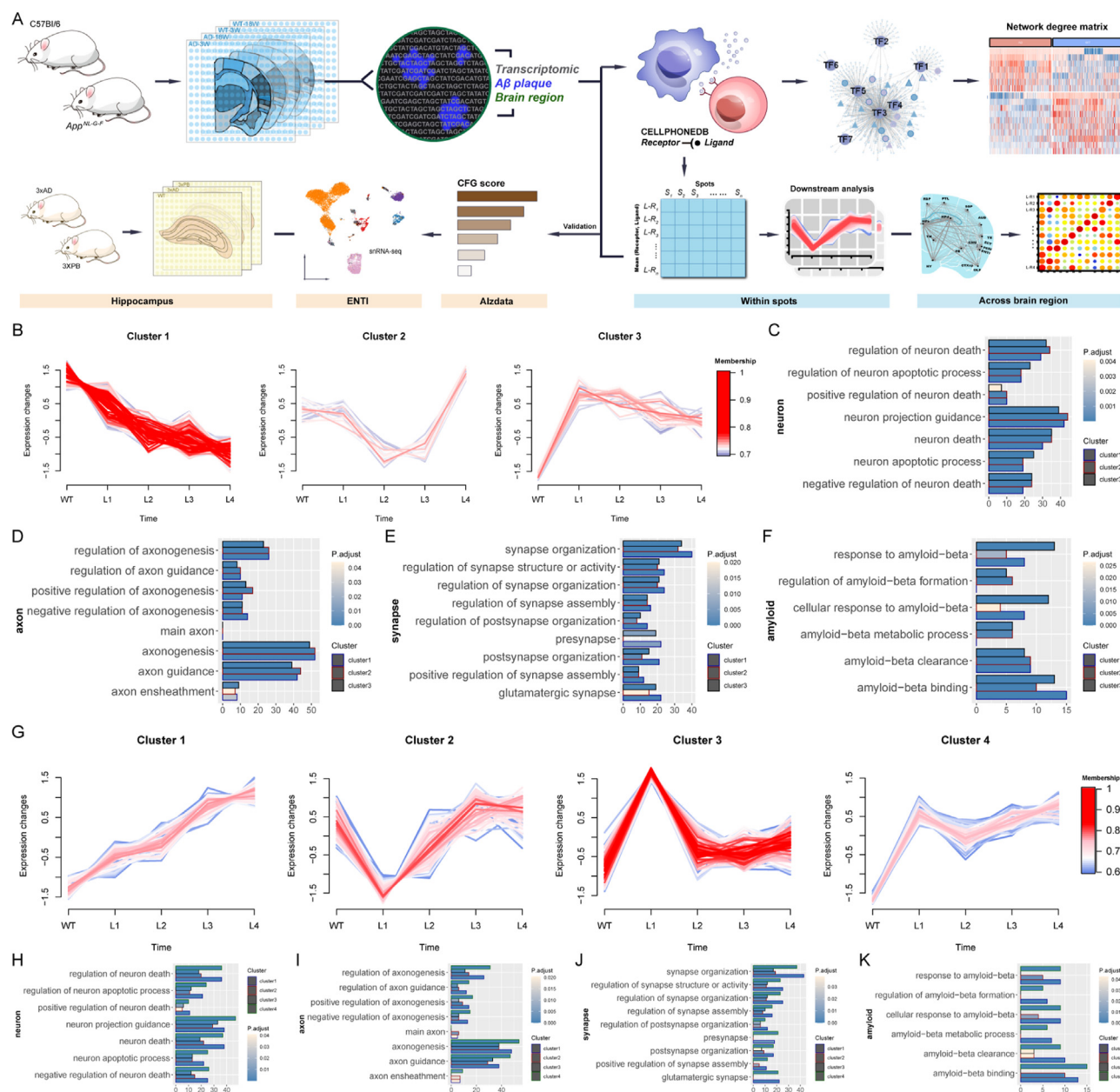


Figure 1 L-R pair clustering and enrichment analysis. **(A)** Workflow of the research. **(B)** Dynamic changes of three L-R clusters in the 3-month spots through Aβ accumulation. The X-axis represents the expression changes and the Y-axis represents different Aβ accumulation groups. Each line represents one L-R pair. The line color represents the membership score for each L-R pair from low (blue) to high (red). **(C–F)** Neuron (C), axon (D), synapse (E), and amyloid (F) related GO analysis results of L-R genes of three L-R clusters in 3 months. The outline color represents different clusters. The fill color represents adjusted P-values from low (blue) to high (yellow). **(G)** Dynamic changes of four L-R clusters in the 18-month spots through Aβ accumulation. The X-axis represents the expression changes and the Y-axis represents different Aβ accumulation groups. Each line represents one L-R pair. The line color represents the membership score for each L-R pair from low (blue) to high (red). **(H–K)** Neuron (H), axon (I), synapse (J), and amyloid (K) related GO analysis results of L-R genes of four L-R clusters in 18 months. The outline color represents different clusters. The fill color represents adjusted P-values from low (blue) to high (yellow).

construct gene networks and scored regulon (gene regulatory modules) activities.⁶ We used GENIE3 in Bioconductor to identify co-expressed gene modules and infer potential TF targets for each module with significantly enriched motifs. RcisTarget identifies enriched transcription factor binding motifs and candidate TFs for a gene list. The motifs that are annotated to the corresponding TF and obtain a

normalized enrichment score >3.0 are retained. Regulons were identified from co-expression and DNA motif analyses. Then, the AUCell function calculated an area under the curve across the ranking of all genes in a particular spot. By calculating whether the input gene subset (TFs and their putative targets in a particular regulon) was enriched at the top of the ranking, AUCell scored the regulon activity in

each spot as a whole, instead of only the TF or individual genes. Regulatory network of high confident annotation interactions (the direct TF-target gene interactions annotated by the cisTarget databases, highConfAnnot = "True") were visualized in Cytoscape (v3.9.1).

Protein-protein interaction network

The protein-protein interaction network was constructed by the Search Tool for the Retrieval of Interacting Genes (STRING) database. The String database provides information about known and predicted protein interactions based on correlation, regulation relationships, or protein binding validated in Co-IP or other experiments. The Cytoscape software v.3.9.1 was used to construct the protein-protein interaction network based on STRING analysis results.

SSN analysis

We performed SSN analysis in Matlab software as previously described.¹⁷ Briefly, the SSN method calculates a statistic $\hat{\rho}_{xy}^{(k)}$ to assess the inter-relationships among gene x and gene y in spot k. "n" was the total spot number. We set $n_x^{(k)} = n_y^{(k)} = 0.1n$ as default in the scatter plot.¹⁷ The coefficient 0.1 denotes the box size. After hypothesis testing based on $\hat{\rho}_{xy}^{(k)}$ to determine the significance of each gene interaction, the total number of significant interactions including gene x is returned as NDM value for gene x. We then used the Wilcoxon rank-sum test to identify genes with significantly different NDM degrees (false discovery rate <0.05) in WT phenotype compared with AD, in different brain regions and age groups.

Results

CellPhoneDB analysis reveals neural related intra-spot L-R interactions

To explore the intra-spot molecular interactions, we clustered L-R pairs by different tendencies through the A β accumulation process. All ST tissue domains were separated into WT and four quantiles of AD according to the A β index (increasing A β accumulation level from L1 to L4; Table S1). All complex information involved in L-R pairs is provided in Table S2. There were three clusters (Fig. 1B and Table S3) in the 3-month group. Cluster 1 showed a downward tendency from WT to L4. Cluster 2 showed an obvious up-regulation from L3 to L4. Compared with any other spot with A β accumulation (L1–L4), L-R pairs in cluster 3 were lower expressed in WT spots. GO analysis revealed the different enrichment results of neuron, axon, synapse, and amyloid related terms in each cluster of the 3-month group (Fig. 1C–F). There are four clusters (Fig. 1G and Table S3) in the 18-month group. Cluster 1 showed an upward tendency from WT to L4. Cluster 2 showed an obvious down-regulation in L1, while cluster 3 was mainly up-regulated in L1. Genes in cluster 4 L-R pairs were expressed at lower levels in WT spots than any other spots with A β accumulation (L1–L4). GO analysis revealed the different enrichment results of neuron, axon, synapse, and amyloid related

terms in each cluster of the 18-month group (Fig. 1H–K). We explored L-R pairs with both genes involved in the same kind of term related to neuron, axon, synapse, or amyloid and found that cluster 3 in the 3-month group and cluster 1 in the 18-month group have the most L-R pairs related to amyloid (Fig. S2). These results showed the dynamic changes of neural related intra-spot L-R interactions through the A β accumulation process.

17 L-R pairs showed opposite tendencies through A β accumulation in the 3-month and 18-month groups

Cluster 1 in the 3-month group and cluster 1 in the 18-month group had 17 common L-R pairs (Table 1). However, these pairs showed opposite tendencies with a general inhibition from WT, L1/L2 (L1 and L2 A β index group), to L3/L4 (L3 and L4 A β index group) in 3 months and a general activation in 18 months among different brain regions (Fig. 1B, G; Fig. 2A). The heatmap showed a similar clustering result of these L-R pairs (except for EPHB2_EFNB2; Fig. 2B) in different brain regions. These results suggested that the different active patterns of 17 common L-R pairs were mainly related to A β accumulation rather than to different brain regions. EPHA3_EFNA3, EPHA1_EFNA3, and EPHA8_EFNA3 have high correlation scores with each other in the 3-month and 18-month groups, respectively, which revealed the existence of potential functional or regulatory interactions between them (Fig. 2C–E).

Next, we performed protein-protein interaction analysis to further explore the interactions in these 17 L-R pairs (Fig. 2F). In this network, 11 genes from EFNA/EFNB and erythropoietin-producing hepatocellular (EPH) gene families as L-R pairs showed strong connections between each other. Their function was mainly related to neuron, axon, and synapse (Fig. S2). Average expression levels of 9 EFNA/

Table 1 List of 17 L-R pairs that showed opposite tendencies through the A β accumulation process in the 3-month and 18-month groups.

L-R pair	Complex
C3_aMb2	aMb2: ITGB2/ITGAM
EFNA1_EPHA8	
FZD9_WNT1	
SPP1_a4b1	a4b1: ITGB1/ITGA4
INS_INSR	
EPHB4_EFNB1	
TNF_PTPRS	
EPHA4_EFNB1	
EFNB2_EPHA4	
ADGRL1_NRG1	
NTF3_NTRK3	
EPHB6_EFNB2	
NRP1_VEGFA	
EPHB2_EFNB2	
EPHA3_EFNA3	
EPHA1_EFNA3	
EPHA8_EFNA3	

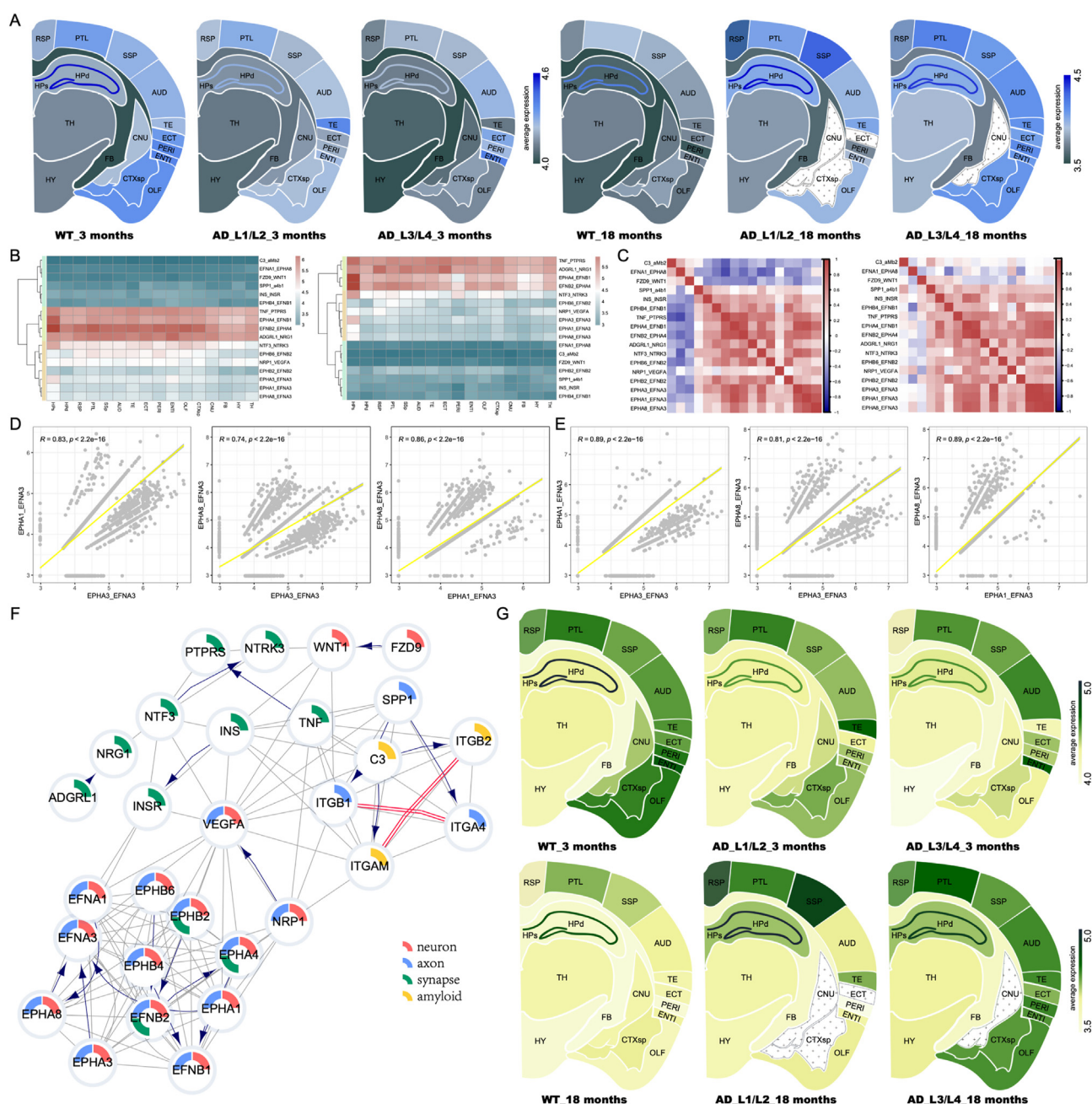


Figure 2 L-R pairs showed opposite tendencies through A β accumulation at 3 and 18 months in intra-spot CellPhoneDB analysis. (A) Average expression of 17 L-R pairs with opposite tendencies. Labels showed sample group information. (B) Expression heatmap of L-R pairs with opposite tendencies in different brain regions at 3 (left) and 18 (right) months. (C) Expression correlation between L-R pairs with opposite tendencies at 3 (left) and 18 (right) months. (D) Correlation between EPHA3_EFNA3, EPHA1_EFNA3, and EPHA8_EFNA3 at 3 months. (E) Correlation between EPHA3_EFNA3, EPHA1_EFNA3, and EPHA8_EFNA3 at 18 months. (F) The protein-protein interaction network of 17 L-R pairs with opposite tendencies. Arc color represents four kinds of GO terms. The directed edge is from ligand to receptor. The red double-line marked edge represents complex. (G) Average expression of EPH-related L-R pairs with opposite tendencies. Labels showed sample group information. TH, thalamus; HY, hypothalamus; FB, fiber tract; HPd, dendritic layer of the hippocampus; HPs, somatic layer of the hippocampus; CNU, cerebral nucleus; CTXsp, cortical subplate; OLF, olfactory area; ENT1, entorhinal area; TE, temporal association area, entorhinal area, and perirhinal area; AUD, auditory area; SSp, primary somatosensory area; PTL, posterior parietal association area; RSP, retrosplenial area.

EPFB/EPH-related L-R pairs were concordant with opposite tendencies that the 3-month group showed an inhibition and the 18-month group showed an activation when progressing from WT, L1/L2, to L3/L4 (Fig. 2G).

C3_ITGAM/ITGB2 complex was the only L-R pair that was involved in amyloid related terms in the protein-protein interaction network. ITGAM was a highly expressed marker gene of microglia (Fig. 3A; single-cell RNA-seq result of different cell types from the human brain in AlzData¹³ (<http://www.alzdata.org/>, GSE67835)). CD11b encoded by ITGAM and CD18 encoded by ITGB2 formed leukocyte-specific complement receptor 3. iC3b is the activation product of complement C3. Complement receptor 3's space conformation will change when interacting with iC3b to initiate A β clearance, phagocytosis, adhesion, or other biological processes¹⁸ (Fig. 3B). C3_ITGAM/ITGB2 complex pair was mainly down-regulated at L3/L4 in the 3-month group, and conversely up-regulated at L3/L4 in the 18-month group (Fig. 3C). These results showed that the C3_ITGAM/ITGB2 complex L-R pair has different correlations with A β index at different ages.

Dynamic changes of L-R interactions across HP

L-R interaction networks of HP showed unique features in different age groups or A β levels (Fig. 4A, B). The L-R interactions in 3 months and 18 months of both WT and AD groups were both most concentrated on the granule cell layer of the dentate gyrus in Hps (HPs_DG_sp) related region pairs. WT spots in both the 3-month and 18-month groups have more significant interactions across HPs_DG_sp and the pyramidal layer of CA3 in Hps (HPs_CA3_sp) compared with L1/L2 and L3/L4 spots in the same age group. Interactions within the DG polymorph layer in HPd

(HPd_DG_po) were more specific in the 18-month spots of the WT group compared with the 3-month spots of the same phenotype group. L3/L4 spots in 3 months showed a general increase in differentially expressed L-R pair numbers compared with WT and L1/L2 spots at this age, especially in HPs_DG_sp itself and HPs_DG_sp to other HP regions. This tendency does not fit in the 18-month group as more region pairs showed less differentially expressed L-R numbers in L3/L4 spots compared with WT and L1/L2 spots. Dot plots showed the top 10 unique L-R pairs (by *P*-values) that were only significantly differentially expressed in one region pair (Fig. 4C). To explore these gene member's functions, we used the AlzData database to search their scores in AD related terms. FLT1 in the 18-month group demonstrated the highest CFG score (5). CXCL13, C3, FGFR2, and CNR2 in the 3-month group and WNT4, VCAM1, and VEGFA in the 18-month group also showed 4 in the CFG grading (Table 2). These results revealed the potential of unique L-R pairs' AD related functions in different HP layers.

Next, we further validated our HP L-R interaction results using the additional spatial transcriptome datasets of two other types of AD model mice. L-R interaction networks of HP showed unique features in different phenotypes (WT, 3xAD, and 3xPB). Compared with 3xAD and 3xPB mice, WT spots had more significant interactions within DG and CA1 subregions and cross CA1/DG (Fig. 5A, B). On the other hand, WT spots had less significant interactions within CA2-3 and cross CA1/CA2-3 compared with spots of 3xAD and 3xPB mice (Fig. 5A, B). Dot plots showed the top 10 unique L-R pairs (by *P*-values) that were only significantly differentially expressed in one region pair within a certain phenotype group (Fig. S3). Next, all unique L-R pairs in 3xAD and 3xPB (11–13 months) HP spots were compared with the first dataset (AppNL-G-F knock-in mice) in L1/L2 and L3/L4 HP spots at 18 months

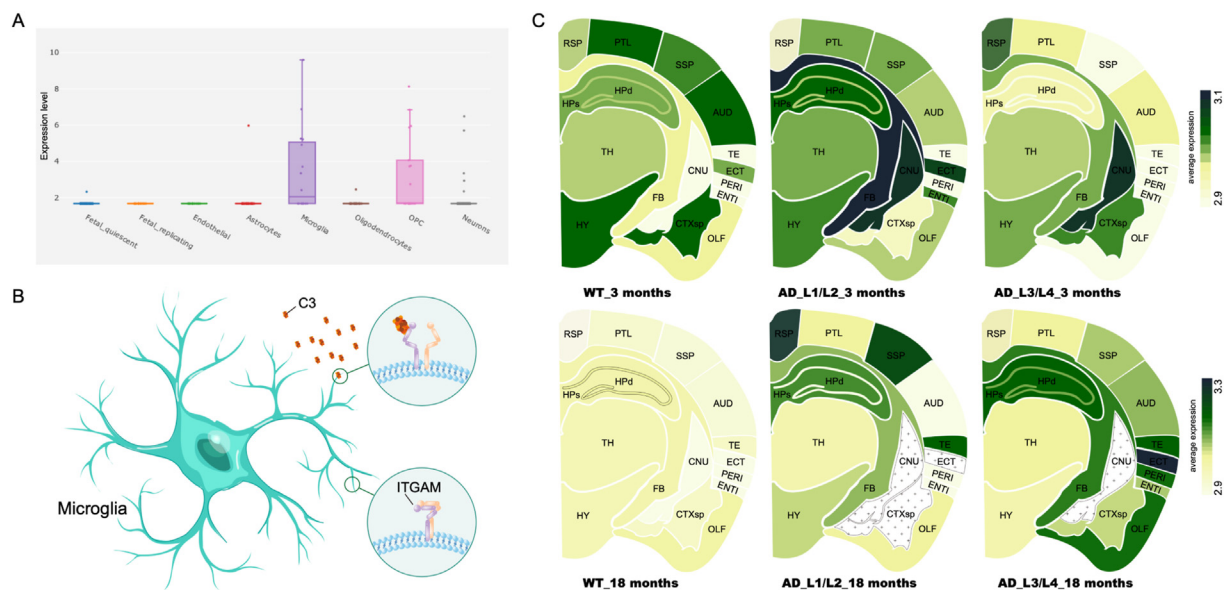


Figure 3 Active pattern of C3-ITGAM/ITGB2 complex. (A) ITGAM expression in different nerve cell types, from the AlzData database. (B) Pattern graph of ITGAM/ITGB2 complex's conformation changes when interacting with C3. (C) Average expression of C3-ITGAM/ITGB2 complex pairs at 3 (top) and 18 (bottom) months. Labels showed sample group information.

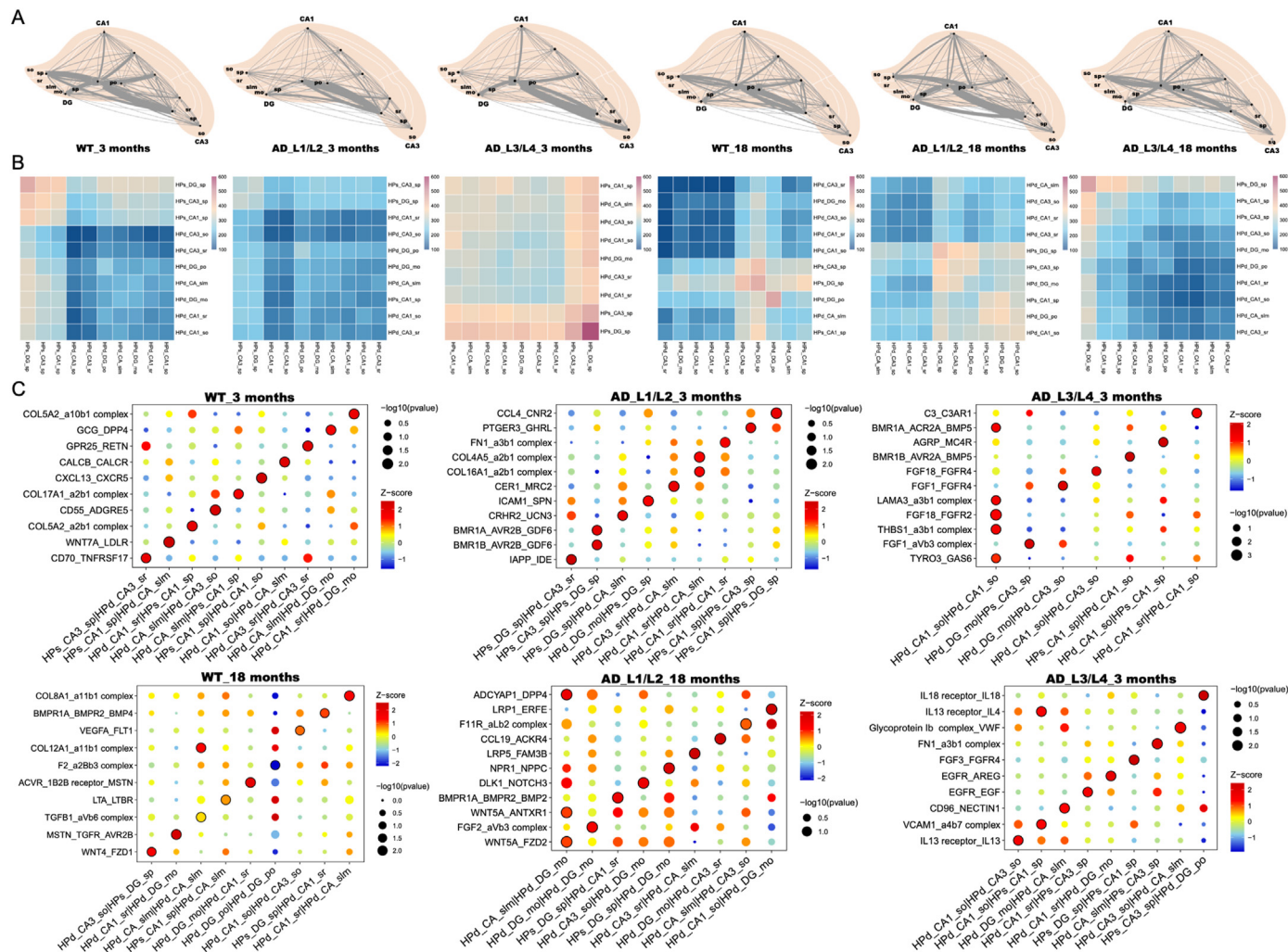


Figure 4 L-R interactions in cross the hippocampal regions in CellPhoneDB analysis. **(A)** L-R interaction networks in different age groups or Aβ levels. The width of the edge represents the number of significantly differentially expressed L-R pairs. The edge direction is from ligand to receptor. Labels showed sample group information. **(B)** Heatmap of significantly differentially expressed L-R pair numbers between two hippocampal layers. **(C)** Unique top 10 L-R pairs (by P-values) that are only significantly differentially expressed in one region pair, marked by dots with black outlines. Labels showed sample group information.

Table 2 CFG scores of top 10 unique L-R pair genes in hippocampus analysis.

Gene	eQTL	GWAS	PPI	Early_DEG	Abeta	Tau	CFG	Note
FLT1	1	2	PSEN1, PSEN2, APOE	Yes	0.466**	−0.119, ns	5	HP18
WNT4	4	2	-	Yes	−0.634***	−0.326, ns	4	HP18
VCAM1	2	0	APOE	Yes	0.575***	0.437, ns	4	HP18
VEGFA	2	0	APP, PSEN2, APOE	Yes	−0.215, ns	−0.758**	4	HP18
CXCL13	1	0	APP, PSEN1, APOE	Yes	0.602***	0.727**	4	HP03
C3	2	0	APP, PSEN1, PSEN2, MAPT	Yes	0.850***	0.761***	4	HP03
FGFR2	3	1	PSEN2, APOE	Yes	0.179, ns	−0.387, ns	4	HP03
CNR2	1	0	APP, PSEN1, APOE	Yes	0.854***	0.750**	4	HP03
ITGA3	1	0	APP, APOE	Yes	−0.284, ns	−0.511, ns	3	HP03, HP18
AREG	NA	0	PSEN2	Yes	0.321*	−0.287, ns	3	HP18
ANTXR1	2	6	-	No	0.535***	−0.003, ns	3	HP18
FN1	1	2	APP, MAPT, APOE	NA	NA	NA	3	HP03, HP18
ITGAV	2	0	APP, PSEN1, MAPT, APOE	Yes	−0.091, ns	0.419, ns	3	HP03, HP18
GP1BA	2	32	APP	NA	NA	NA	3	HP18
IL13RA1	1	NA	-	Yes	0.727***	0.588*	3	HP18
TGFB1	1	0	APP, MAPT, APOE	NA	0.871***	0.681**	3	HP18
EGFR	2	2	PSEN1, PSEN2, APOE	NA	0.119, ns	0.012, ns	3	HP18
F11R	1	0	NA	Yes	0.801***	0.639*	3	HP18
VWF	2	0	APP, MAPT, APOE	NA	0.640***	0.493, ns	3	HP18
FZD2	9	0	APP, PSEN1, MAPT, APOE	NA	−0.419**	−0.702**	3	HP18
COL4A5	4	NA	APOE	No	0.396**	0.359, ns	3	HP03
COL16A1	1	0	-	Yes	0.470**	0.176, ns	3	HP03
FGF1	0	17	PSEN2, APOE	Yes	0.145, ns	−0.415, ns	3	HP03
FGF18	10	1	PSEN2, MAPT, APOE	NA	NA	NA	3	HP03
CCL4	2	0	-	Yes	0.873***	0.887***	3	HP03
CD70	1	0	-	Yes	0.714***	0.691**	3	HP03
WNT7A	1	0	PSEN2	Yes	−0.055, ns	0.241, ns	3	HP03
TNFRSF17	2	0	PSEN2	NA	0.775***	0.574*	3	HP03

Notes: eQTL, expression of the target gene is regulated by AD genetic variants (genetic variants: IGAP GWAS P -value $< 1E-3$; eQTL: P -value $< 1E-3$). GWAS, IGAP P -value $< 1E-3$. PPI, the target gene has significant physical interaction with APP, PSEN1, PSEN2, APOE, or MAPT (P -value < 0.05). Early_DEG, the target gene is differentially expressed in AD mouse models before AD pathology emergence. Pathology cor (abeta), correlation of target gene expression with AD pathology in abeta line AD mouse models (r , P -value; ns: $P > 0.05$; * $P < 0.05$; ** $P < 0.01$; *** $P < 0.001$). Pathology cor (tau), correlation of target gene expression with AD pathology in tau line AD mouse models (r , P -value; ns: $P > 0.05$; * $P < 0.05$; ** $P < 0.01$; *** $P < 0.001$). CFG is the total convergent functional genomic score of the target gene; 1 CFG point is added if any of the above evidence is significant, and the CFG point ranges from 0 to 5.

respectively. Although 3xAD and 3xPB mice were younger than the AppNL-G-F knock-in mice, they were all in the cognitive impairment disease stage with deficits in memory and learning. "COL1A1_a10b1 complex", "COL8A2_a10b1 complex", "NRG3_ERBB4", "MADCAM1_a4b7 complex", and "EPHA3_EFNA3" were significantly differentially expressed in the same region pairs in 3xAD and 3xPB spots. Meanwhile, these L-R pairs were significantly differentially expressed in L1/L2 subregion pairs at 18 months which belong to the same region that the unique L-R pairs in 3xAD and 3xPB spots showed different expression (Fig. 5C). "NAMPT_P2RY6", "COL13A1_a1b1 complex", "MERTK_GAS6", "NPR1_NPPC", and "NRG3_ERBB4" were significantly differentially expressed in the same region pairs in 3xAD and 3xPB spots. Meanwhile, these LR pairs were significantly differentially expressed in L3/L4 subregion pairs at 18 months which belong to the same regions that the unique L-R pairs in 3xAD and 3xPB spots showed different expression (Fig. 5D). These results showed the potential functional interactions in the

related region pairs and further validated the spatial conservation of these L-R pairs in different AD models.

TF regulatory analysis identified a functional TF network in the HP

To further explore other kinds of interactions in HP, we used SCENIC to perform TF regulatory analysis. According to the TF regulatory network activity heatmap, different phenotype groups at 3 months and 6 months were clustered together, whereas the transcriptomics regulatory profile showed a clear separation between WT and AD groups at 12 months and 18 months (Fig. 6A). This separation was concordant with the progression of AD pathology over these periods. According to the TF regulatory activity heatmap of different HP regions, HPs (somatic layer of the hippocampus), and HPd (dendritic layer of the hippocampus) spots were clustered as two groups as expected (Fig. 6B). Regulators were clustered into three groups according to their

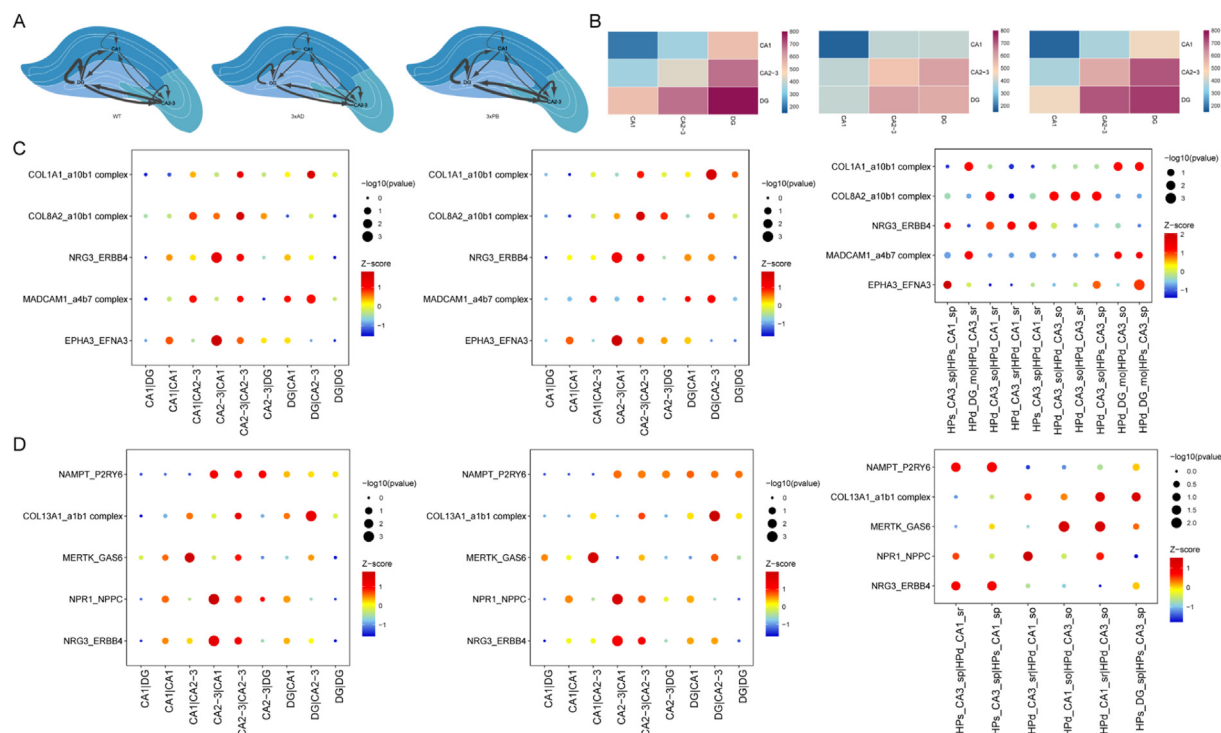


Figure 5 L-R interactions cross the hippocampus in 3xAD and 3XPB mice. (A) L-R interaction networks in WT, 3xAD, and 3XPB mice. The width of the edge represents the number of significantly differentially expressed L-R pairs. The edge direction is from ligand to receptor. (B) Heatmap of significantly differentially expressed L-R pair numbers between two hippocampal layers. Different colors represented the numbers of differentially expressed L-R pairs from less (blue) to more (red). (C) Significantly differentially expressed L-R interactions in the same region pairs in 3xAD, 3XPB slides, and corresponding subregion pairs in the L1/L2 group at 18 months. (D) Significantly differentially expressed L-R interactions in the same region pairs in 3xAD, 3XPB slides, and corresponding subregion pairs in the L3/L4 group at 18 months.

activities. Cluster 1 regulons were mainly active in the somatic layer, whereas cluster 2 regulons were mainly active in the DG_{po} subregion of the dendritic layer. Genes in cluster 1 regulons were involved in neurotransmitter (regulation of neurotransmitter levels, neurotransmitter transport, neurotransmitter secretion, etc.), synaptic (synaptic vesicle cycle, synapse organization, synaptic vesicle transport, etc.), and cognition/memory related functions (Fig. 6C). Genes in cluster 2 regulons were involved in stimulus (especially visual stimulus) response related functions like visual perception, detection of light stimulus, etc. (Fig. 6D). Further, we used Cytoscape for network visualization and marked the categories of gene function, i.e., cognition/memory, neurotransmitter, and synaptic in cluster 1 regulon network (Fig. 6E) and stimulus response in cluster 2 regulon network (Fig. 6F).

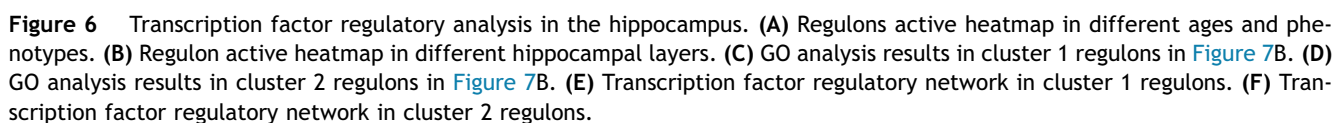
In the TF regulatory heatmap of different A β index groups, we found a cluster (cluster A) that included 6 up-regulated regulons from WT-to-L4 A β levels (Fig. 7A). GO analysis showed that their functions were mainly involved in "negative regulation of humoral immune response" and "DNA-binding transcription activator activity, RNA polymerase II-specific" (Fig. 7B). Different gene functions were also marked in the regulatory network of cluster A (Fig. 7C). *PRKCB* and *NPTX2* in HP network and *NR2F2* in cluster A demonstrated the highest CFG score, suggesting these

genes may be crucial in AD related pathological process (Table 3).

TF regulatory analysis identified functional TF network in ENTI

Since ENTI is an important affected region in the early AD process,^{10,11} we further performed SCENIC analysis in this brain area. In the TF regulatory heatmap of different A β index groups, we found cluster 3 with 32 regulons was mainly active in L1 and L2 groups (Fig. 7D). This result was concordant with the progression of AD pathology in this region. GO analysis showed functions of regulons in cluster 3 were mainly involved in neurotransmitter (regulation of neurotransmitter levels, neurotransmitter transport, etc.), synaptic (synapse organization, synaptic vesicle cycle, synaptic vesicle localization, etc.), and cognition/memory related terms (Fig. 7E). Regulatory network of cluster 3 was marked with the corresponding gene functions (Fig. 7F). *MEF2C* and *PRKCB* ranked first in the CFG score list of cluster 3 TFs (Table 4). These findings suggest that *MEF2C* and *PRKCB* may potentially regulate the neuropathology process in ENTI.

To determine if any of the high CFG score genes related to ENTI were differentially expressed in human AD



SSN analysis revealed specific high NDM degree genes in different phenotypes

To explore gene connections at the whole transcriptome level, we constructed SSN for each single spot (*i.e.*, one network for one spot) in HP and ENTI regions. Gene expression changes dynamically or even drastically with time and conditions, however, the functions or linear/nonlinear associations among genes generally remain unchanged or change gradually with small perturbations.¹⁷ So, we used SSN to transform the “unstable” gene expression matrix into a “stable” gene association based NDM. We

Table 3 CFG scores of target genes in hippocampal transcription factor analysis.

Gene	eQTL	GWAS	PPI	Early_DEG	abeta	tau	CFG	Cluster
PRKCB	1	5	PSEN1, MAPT, APOE	Yes	-0.231, ns	-0.199, ns	4	HPs
NR2F2	0	1	PSEN2, APOE	Yes	-0.487***	0.075, ns	4	A
NPTX2	1	1	-	Yes	-0.688***	-0.783***	4	HPs
SYT1	0	7	MAPT	Yes	-0.209, ns	-0.471, ns	3	HPs
SV2A	2	0	NA	Yes	-0.199, ns	-0.746**	3	HPs
STAT5A	1	0	PSEN2	NA	0.477**	0.606*	3	A, HPd
SREBF2	1	2	APOE	NA	NA	NA	3	HPs
HSF1	1	0	-	Yes	0.462**	0.120, ns	3	HPs
HOXA5	1	NA	-	Yes	0.372*	0.052, ns	3	HPs
HOMER1	1	0	-	Yes	-0.338*	-0.341, ns	3	HPs
GRIA1	3	30	MAPT	No	0.036, ns	0.145, ns	3	HPs
GAD1	3	0	-	Yes	-0.207, ns	-0.562*	3	HPs
ESRRA	2	0	PSEN2, MAPT, APOE	Yes	-0.043, ns	-0.309, ns	3	A, HPs
EGR1	0	2	PSEN2, MAPT	NA	-0.403**	-0.559*	3	HPs
ARC	2	1	-	Yes	-0.194, ns	-0.259, ns	3	HPs

Notes: eQTL, expression of the target gene is regulated by AD genetic variants (genetic variants: IGAP GWAS P -value $< 1E-3$; eQTL: P -value $< 1E-3$). GWAS, IGAP P -value $< 1E-3$. PPI, the target gene has significant physical interaction with APP, PSEN1, PSEN2, APOE, or MAPT (P -value < 0.05). Early_DEG, the target gene is differentially expressed in AD mouse models before AD pathology emergence. Pathology cor (abeta), correlation of target gene expression with AD pathology in abeta line AD mouse models (r , P -value; ns: $P > 0.05$; * $P < 0.05$; ** $P < 0.01$; *** $P < 0.001$). Pathology cor (tau), correlation of target gene expression with AD pathology in tau line AD mouse models (r , P -value; ns: $P > 0.05$; * $P < 0.05$; ** $P < 0.01$; *** $P < 0.001$). CFG is the total convergent functional genomic score of the target gene; 1 CFG point is added if any of the above evidence is significant, and the CFG point ranges from 0 to 5.

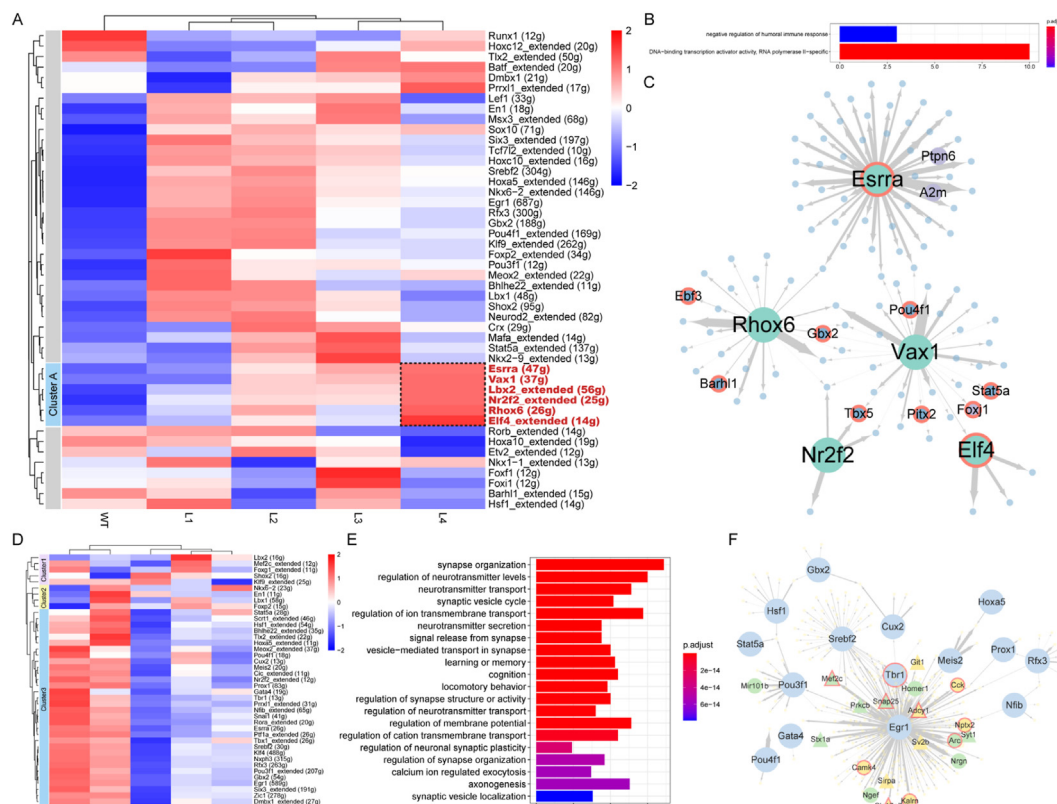


Figure 7 Transcription factor regulatory analysis in A β index group. (A) Regulon active heatmap in different A β index groups in the hippocampus. (B) GO analysis results in cluster A regulons in Figure 8A. (C) Transcription factor regulatory network in cluster A regulons. (D) Regulon active heatmap in different A β index groups in the entorhinal cortex. (E) GO analysis results in cluster 3 regulons in Figure 7D. (F) Transcription factor regulatory network in cluster 3 regulons.

Table 4 CFG scores of target genes in entorhinal cortex transcription factor analysis.

Gene	eQTL	GWAS	PPI	Early_DEG	abeta	tau	CFG
MEF2C	1	2	PSEN2, MAPT	Yes	-0.159, ns	-0.198, ns	4
PRKCB	1	5	PSEN1, MAPT, APOE	Yes	-0.231, ns	-0.199, ns	4
SV2A	2	0	NA	Yes	-0.199, ns	-0.746**	3
VGF	2	0	-	Yes	-0.792***	-0.760**	3
HSF1	1	0	-	Yes	0.462**	0.120, ns	3
SREBF2	1	2	APOE	NA	NA	NA	3
EGR1	0	2	PSEN2, MAPT	NA	-0.403**	-0.559*	3
STAT5A	1	0	PSEN2	NA	0.477**	0.606*	3
CUX2	2	2	PSEN2	NA	NA	NA	3
GRIA1	3	30	MAPT	No	0.036, ns	0.145, ns	3

Notes: eQTL, expression of the target gene is regulated by AD genetic variants (genetic variants: IGAP GWAS P -value $< 1E-3$; eQTL: P -value $< 1E-3$). GWAS, IGAP P -value $< 1E-3$. PPI, the target gene has significant physical interaction with APP, PSEN1, PSEN2, APOE, or MAPT ($P < 0.05$). Early_DEG, the target gene is differentially expressed in AD mouse models before AD pathology emergence. Pathology cor (abeta), correlation of target gene expression with AD pathology in abeta line AD mouse models (r , P -value; ns: $P > 0.05$; $*P < 0.05$; $**P < 0.01$; $***P < 0.001$). Pathology cor (tau), correlation of target gene expression with AD pathology in tau line AD mouse models (r , P -value; ns: $P > 0.05$; $*P < 0.05$; $**P < 0.01$; $***P < 0.001$). CFG is the total convergent functional genomic score of the target gene; 1 CFG point is added if any of the above evidence is significant, and the CFG point ranges from 0 to 5.

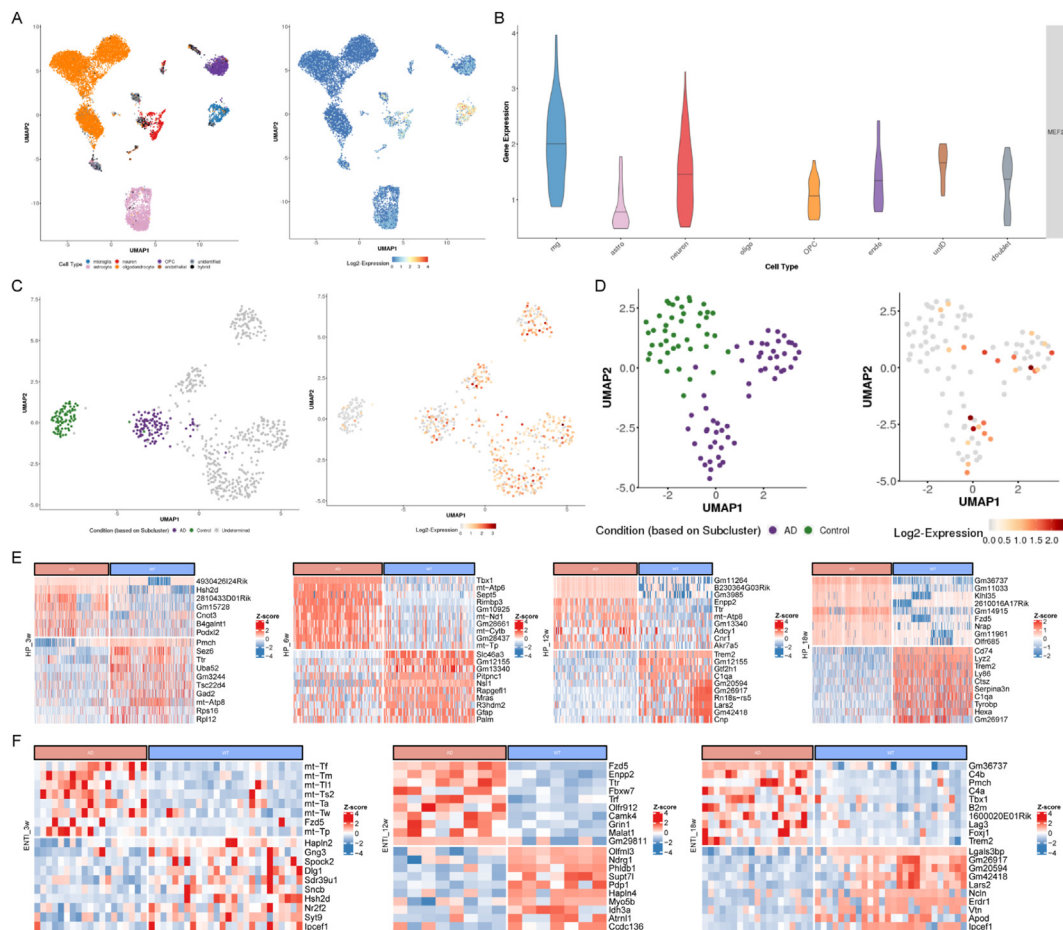


Figure 8 snRNA-seq in human ENT1 and SSN analysis results. (A) UMAP manifold of snRNA-seq colored by cell type (left) and MEF2C expression (right). (B) MEF2C expression in different cell types. (C) UMAP of neuron subset colored by phenotype (left) and MEF2C expression (right). (D) UMAP of endothelial cell subset colored by phenotype (left) and MEF2C expression (right). (E) Genes with significantly different NDM degrees (WT vs. AD) in the hippocampus. (F) Genes with significantly different NDM degrees (WT vs. AD) in ENT1. snRNA-seq, single-nuclei RNA sequencing; ENT1, entorhinal cortex; SSN, spot-specific network; WT, wild-type; AD, Alzheimer's disease; NDM, network degree matrix.

found genes with significant differences in NDM degrees between WT and AD in HP (Fig. 8E) and ENT1 (Fig. 8F) in different age groups. For example, *Ttr* showed a higher NDM degree in WT spots than AD spots in the 3-month group of HP. Conversely, *Ttr* showed a higher NDM degree in AD spots than WT spots in the 12-month group of HP. *Trem2* showed a higher NDM degree in WT spots than AD spots in the 18-month group of HP. Conversely, *Trem2* showed a higher NDM degree in AD spots than WT in the 18-month group of ENT1. These results showed the specific high NDM degree genes that may have wide interactions across the whole transcriptomes in certain phenotypes in different brain regions and age groups.

Discussion

In the current study, we performed L-R communication, TF regulatory network, and SSN analyses to reveal the dependence and dynamics of gene associations from multiple perspectives based on samples from mouse and human brains. Using an ST dataset of *App* knock-in mouse model,¹² we revealed the specific L-R interaction status across the whole brain areas or HP layers within and across spots in different spatial regions and pathological status. We also further revealed important TF regulatory and SSN gene associations in HP and ENT1, respectively. Next, we used another independent ST dataset from different AD mouse models and human ENT1 snRNA-seq data and AlzData database (that includes both human and mouse data) to partially validate our results. This study provides a framework for the comprehensive analyses of gene-gene associations or transcriptional networks that are spatio-temporal dependent within or among spots in ST, thereby laying the foundation for exploring characteristic gene functions and in-depth AD etiology from the perspective of spatio-temporal dependent gene interactions.

In our study, Mfuzz assigned neuron, axon, synapse, and amyloid related intra-spot L-R pairs into different clusters. Interestingly, we found that 17 L-R pairs showed opposite tendencies in the 3-month and 18-month groups through the A β accumulation process. The opposite tendencies of these 17 L-R pairs may be related to their different functions in different pathological statuses. For example, interactions in the protein-protein interaction network were most concentrated among EFNA/EFNB and EPH gene families. EPH receptors belong to the tyrosine kinase receptor gene family that plays a pivotal role in brain development.¹⁹ EPHA4 activation can lead to dendritic spine retraction and synaptic dysfunction.¹⁹ Several reports also support the protective role of EPHA4 in AD pathogenesis.^{20,21} EPHA4 promotes the proliferation of neural progenitor cells and alleviates cognitive impairment in a PDGFR β dependent manner.²⁰ These studies showed the contradictory functions of EPHA4 in AD pathogenesis. Since our results also showed opposite tendencies in EPHA4 related L-R pairs through A β accumulation in the 3-month and 18-month groups, further investigation may focus on the potentially different functions of EPHA4 in the different stages or processes of the disease during different age groups.

C3_ITGAM/ITGB2 complex was the only A β related L-R pair between spots in the GO enrichment analysis results

that has the opposite tendencies between 3-month-old and 18-month-old mice. Interaction of complement component iC3b with its receptor complement receptor 3 (CD11b/CD18) on the surface of microglia involved in the uptake and clearance of A β .²² C3 deficiency in APP/PS1 mice increased plaque burden in the mouse brain with decreased age- and plaque-related synapse and neuron loss, glial reactivity, and spared cognitive decline.¹⁸ Our results showed the potential of the C3_ITGAM/ITGB2 complex's different functions in plaque-related neurodegeneration and cognitive health at different age groups.

L-R associations across brain regions showed different states between different ages, region pairs, and phenotypes in both expression levels and different numbers of significantly differentially expressed L-R pairs. For example, VCAM1_a4b7 complex in HPd_CA1_so to HPs_CA1_sp, L3/L4 level in 18 months. The low levels of VCAM1 on brain endothelial cells allowed microglia to remain ramified, calm, and proceed to neurogenesis. Increased VCAM1 expression in the old brain tethered more leukocytes to the luminal side and activated microglia to suppress neurogenesis which could be reversed by systemic anti-VCAM1 antibody.²³ In addition, some L-R pairs showed spatial conservative differential expression (differentially expressed in the same region pair when compared with other region pairs) in different AD models (*App*^{NL-G-F}, 3xAD, and 3xPB) including collagen related genes like *COL1A1*, *COL8A2*, and *COL13A1*. Post-mortem evidences were reported in transgenic animal models and brains of individuals with AD, that increased collagen content was related to microvascular morphology changes along with the cognitive status worsening in AD mouse model and AD human patients.^{1,24} Our results supplied the spatio-temporal information for the interactions that contained these AD related genes. These results may furnish a foundation for future functional research based on L-R interactions instead of only individual genes, with the novel high spatial resolution empowered by ST.

According to the TF regulatory network analysis results of HP in different age groups, transcriptomics regulatory profiles of WT mice at 12 and 18 months of age were clustered together, whereas the AD mice transcriptomics regulatory profiles still changed between these two time points. These tendencies were in concordance with the progression of pathology over these periods, as well as the original research results of this dataset.¹² GO enrichment analysis results showed that regulons active in the somatic layer were involved in neurotransmitter, synaptic, and cognition/memory functions, whereas regulons active in the DG_po subregion of the dendritic layer were involved in stimulus (e.g., visual stimulus) response related functions. These results showed region-specific active patterns of such regulons in HP with different potential functions.

Next, we explored the regulon active patterns through A β deposition in HP and found that a gradually up-regulated regulon cluster (cluster A) was involved in "negative regulation of humoral immune response" and "DNA-binding transcription activator activity, RNA polymerase II-specific" GO terms. These dynamic changes showed cluster A regulons may be related to the humoral immune response and the catastrophic failure of the transport of transcription-related molecules (especially RNA polymerase II) between the cytoplasm and the nucleus in the late AD neurotoxic

process.²⁵ GO analysis results of 32 active regulons in L1 and L2 groups of ENT1, an early affected area in AD, revealed their wide influences at the early AD stage including neurotransmitter, synaptic, and cognition/memory related functions. *MEF2C* was one of the high CFG score TFs in these active regulons. *MEF2C* was highly expressed in AD neuron and endothelial cell clusters compared with control neuron and endothelial cell clusters in the snRNA-seq data of human ENT1 (Fig. 8C, D). The *MEF2* TF family modulates the structural and synaptic plasticity underlying memory formation, and *MEF2C* gene is related to synaptic function alteration in AD.^{26–29} Overexpression of *MEF2* has been reported to impede the increase of learning-induced spine and impair memory formation via an Arc-mediated reduction of GluA2-AMPA-type glutamate receptor expression.^{30,31}

Gene connection analysis at the whole transcriptome level showed genes with high NDM degrees were specific in certain phenotypes of different regions and age groups. *Ttr* is an AD neuroprotection gene that is known to bind to A β and facilitate its clearance from the brain,³² SSN analysis showed the transcriptome-based influences of *Ttr* were unique in certain phenotypes in different age groups, which underlying the A β clearance function of *Ttr* may also depend on age conditions. *Trem2* is an AD risk factor due to its impacts on central/peripheral lipid metabolism and the integrity of the blood-brain barrier.³³ SSN analysis showed the transcriptome-based influences of *Trem2* were unique in certain phenotypes in different brain areas. These results showed these genes' unique transcriptome-based influences in different phenotypes, regions, and age groups that further suggested their known impact on AD may be unique under such different conditions. Here, this added information was explored at the novel ST resolution for the first time.

CellPhoneDB and SCENIC were developed for the single-cell RNA-seq data. Both methods have been used in the bulk-like ST data before.^{34–36} However, the related results were limited to the spot resolution, without specified cell type information.

In the case of the validation in the HP 3xAD/3xPB datasets, we tried to identify the L-R interactions that were significantly differentially expressed in the same region pairs in 3xAD, 3xPB slides, and the same corresponding subregion pairs in L1/L2 or L3/L4 (Fig. 4C) group at 18 months of the AppNL-G-F dataset, respectively. These results primarily validated that the significantly differentially expressed L-R interactions were consistent in different AD mouse models. In the case of the validation in the human ENT1 dataset, we primarily validated that some important TFs in the AppNL-G-F dataset were also differentially expressed in certain cell types of human AD patients compared with control (Fig. 8A–D). However, a limitation of the present study is the low resolution of the ST dataset. The research object of this study is the A β niche (based on spots). Both cell proportion and regulon activity of certain cell types can affect the regulon activity level of spots. Under the current resolution of this dataset, it is more suitable for conducting preliminary research in the A β niche (100 μ m in diameter), rather than excessive distinguishing between the two reasons mentioned above. A higher-resolution ST dataset and related molecular experiments will

provide more thoroughness and more solid validation of the current results.

Conclusions

The current study revealed gene associations in multi-aspects based on an ST dataset of *App* knock-in mouse model with the validation on the Alzdata database and supplementary spatial sequencing/snRNA-seq datasets from mice and human brains. The unique genome-wide spatio-temporal dependent association features in different spatial, temporal, and pathological statuses establish the foundations to reveal the advanced and in-depth AD etiology and mechanisms in different stages of the neurodegeneration processes on the amyloid plaque niche in AD.

Author contributions

SW, HS, and HD conceived and designed the study. SW analyzed data and wrote the manuscript. JG, CQ, RS, MP, YG, HS, HX, and HD revised the manuscript. HX contributed to funding acquisition. All authors read and approved the manuscript.

Conflict of interests

The authors declare no conflict of interests.

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Data availability

The whole brain ST dataset can be accessed with accession number GSE152506 in GEO database. The HP ST dataset was available at Mendeley <https://doi.org/10.17632/6s959w2zyr.1y>. The ENT1 snRNA-seq dataset can be accessed with accession number GSE138852 in GEO database.

Appendix A. Supplementary data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.gendis.2024.101337>.

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