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Integrative Analysis to Investigate Complex Interaction in Alzheimer's Disease

Zeran Li
Washington University in St. Louis

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Division of Biology and Biomedical Sciences
Neurosciences

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Integrative Analysis to Investigate Complex Interaction in Alzheimer’s Disease
by
Zeran Li

A dissertation presented to
The Graduate School
of Washington University in
partial fulfillment of the
requirements for the degree
of Doctor of Philosophy

May 2019
St. Louis, Missouri
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<th>Description</th>
</tr>
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<tbody>
<tr>
<td>Aβ</td>
<td>Amyloid β</td>
</tr>
<tr>
<td>AD</td>
<td>Alzheimer’s Disease</td>
</tr>
<tr>
<td>ADAD</td>
<td>Autosomal Dominant Alzheimer’s Disease</td>
</tr>
<tr>
<td>ALS</td>
<td>Amyotrophic lateral sclerosis</td>
</tr>
<tr>
<td>AMP-AD</td>
<td>Advanced Medicines Partnership - Alzheimer’s Disease</td>
</tr>
<tr>
<td>APC</td>
<td>Anterior prefrontal cortex</td>
</tr>
<tr>
<td>BM9</td>
<td>Dorsal lateral prefrontal cortex</td>
</tr>
<tr>
<td>BM10</td>
<td>Anterior prefrontal cortex</td>
</tr>
<tr>
<td>BM22</td>
<td>Superior temporal gyrus</td>
</tr>
<tr>
<td>BM24</td>
<td>Ventral anterior cingulate cortex</td>
</tr>
<tr>
<td>BM36</td>
<td>Parahippocampal gyrus</td>
</tr>
<tr>
<td>BM44</td>
<td>Inferior frontal gyrus</td>
</tr>
<tr>
<td>CB</td>
<td>Cerebellum</td>
</tr>
<tr>
<td>CDR</td>
<td>Clinical Dementia Rating</td>
</tr>
<tr>
<td>CERAD</td>
<td>The Consortium to Establish a Registry for Alzheimer’s Disease</td>
</tr>
<tr>
<td>cQTL</td>
<td>cell type QTL</td>
</tr>
<tr>
<td>CSF</td>
<td>Cerebrospinal fluid</td>
</tr>
<tr>
<td>CTX</td>
<td>Cortex</td>
</tr>
<tr>
<td>DIAN</td>
<td>Dominantly Inherited Alzheimer Network</td>
</tr>
<tr>
<td>DLB</td>
<td>Dementia with Lewy bodies</td>
</tr>
<tr>
<td>DLPFC</td>
<td>Dorsal lateral prefrontal cortex</td>
</tr>
<tr>
<td>DSA</td>
<td>Digital Sorting Algorithm</td>
</tr>
<tr>
<td>EOAD</td>
<td>Early-onset Alzheimer’s disease</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
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<tr>
<td>--------------</td>
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<tr>
<td>FCX</td>
<td>Frontal cortex</td>
</tr>
<tr>
<td>FTD</td>
<td>Frontotemporal dementia</td>
</tr>
<tr>
<td>FTLD</td>
<td>Frontotemporal lobar degeneration</td>
</tr>
<tr>
<td>GO term</td>
<td>Gene Ontology Enrichment Analysis</td>
</tr>
<tr>
<td>IFG</td>
<td>Inferior frontal gyrus</td>
</tr>
<tr>
<td>IGAP</td>
<td>The International Genomics of Alzheimer’s Project</td>
</tr>
<tr>
<td>iPSC</td>
<td>Induced pluripotent stem cell</td>
</tr>
<tr>
<td>Knight ADRC</td>
<td>Charles F. and Joanne Knight Alzheimer’s Disease Research Center</td>
</tr>
<tr>
<td>KEGG</td>
<td>Kyoto Encyclopedia of Genes and Genomes</td>
</tr>
<tr>
<td>LOAD</td>
<td>Late-onset Alzheimer’s disease</td>
</tr>
<tr>
<td>meanProfile</td>
<td>Implementation of method Population-Specific Expression Analysis</td>
</tr>
<tr>
<td>MSBB</td>
<td>Mount Sinai Brain Bank</td>
</tr>
<tr>
<td>MSSM</td>
<td>Mount Sinai School of Medicine</td>
</tr>
<tr>
<td>NIA</td>
<td>National institute of aging</td>
</tr>
<tr>
<td>NPCs</td>
<td>Neural progenitor cells</td>
</tr>
<tr>
<td>PA</td>
<td>Pathological aging</td>
</tr>
<tr>
<td>PAR</td>
<td>Parietal cortex</td>
</tr>
<tr>
<td>PC</td>
<td>Principal component</td>
</tr>
<tr>
<td>PCA</td>
<td>Principal component analyses</td>
</tr>
<tr>
<td>PD</td>
<td>Parkinson’s disease</td>
</tr>
<tr>
<td>PHG</td>
<td>Parahippocampal gyrus</td>
</tr>
<tr>
<td>PMI</td>
<td>Post-mortem index</td>
</tr>
<tr>
<td>PSEA</td>
<td>Population-Specific Expression Analysis</td>
</tr>
<tr>
<td>PSP</td>
<td>Progressive Supranuclear Palsy</td>
</tr>
<tr>
<td>ptau</td>
<td>Phosphorylated tau</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
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<td>--------------</td>
<td>-------------</td>
</tr>
<tr>
<td>QC</td>
<td>Quality control</td>
</tr>
<tr>
<td>RIN</td>
<td>RNA integrity number</td>
</tr>
<tr>
<td>rmcorr</td>
<td>Repeated measures correlation</td>
</tr>
<tr>
<td>RMSE</td>
<td>Root-mean-square error</td>
</tr>
<tr>
<td>SCZ</td>
<td>Schizophrenia</td>
</tr>
<tr>
<td>SNP</td>
<td>Single nucleotide polymorphism</td>
</tr>
<tr>
<td>ssNMF</td>
<td>Semi-supervised non-negative matrix factorization</td>
</tr>
<tr>
<td>STG</td>
<td>Superior temporal gyrus</td>
</tr>
<tr>
<td>TC</td>
<td>Temporal cortex</td>
</tr>
<tr>
<td>TIN</td>
<td>Transcript integrity number</td>
</tr>
<tr>
<td>TOM</td>
<td>Topological overlap matrix</td>
</tr>
<tr>
<td>TRAP-seq</td>
<td>Translating ribosome affinity purification sequencing</td>
</tr>
<tr>
<td>WGCNA</td>
<td>Weighted correlation network analysis</td>
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</table>
Acknowledgments

Firstly, I would like to express my gratitude to my advisor Dr. Carlos Cruchaga for his support of my PhD study and related research. I thank him for his patience, encouragement, insights, and high standard that lead me through my PhD training. I appreciate all the learning and training opportunities he has provided to me. His guidance and mentorship helped me in all the time of research and writing of this thesis.

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Zeran Li

Washington University in St. Louis

May 2019
Dedicated to my parents.
ABSTRACT OF THE DISSERTATION

Integrative Analysis to Investigate Complex Interaction in Alzheimer’s Disease

by

Zeran Li

Doctor of Philosophy in Biology and Biomedical Sciences

Neurosciences

Washington University in St. Louis, 2019

Carlos Cruchaga, Chair

Alzheimer’s disease (AD) is a neurodegenerative disorder featuring progressive cognitive and functional deficits. Pathologically, AD is characterized by tau and amyloid β protein deposition in the brain. As the sixth leading cause of death in the U.S., the disease course usually last from 7 to 10 years on average before the consequential death. In 2019 there are estimated 5.8 million Americans living with AD affecting 16 million family members. At certain stage of the disease course, patients with inability of maintaining their daily functioning highly depend on caregivers, primarily family caregivers, that incur estimated 18.4 billion unpaid hours of cares, which is equivalent to 232 billion dollars. These huge economic burdens and inevitable emotional distress on the family and the society would also increase as the number of AD affected population could triple by 2050.

Altered cellular composition is associated with AD progression and decline in cognition, such as neuronal loss and astrocytosis, which is a key feature in neurodegeneration but has often been overlooked in transcriptome research. To explore the cellular composition changes in AD, I developed a deconvolution pipeline for bulk RNA-Seq to account for cell type specific effects in brain tissues. I found that neuronal and astrocyte relative proportions differ between healthy and
diseased brains and also among AD cases that carry specific genetic risk variants. Brain carriers of pathogenic mutations in APP, PSEN1, or PSEN2 presented lower neuron and higher astrocyte relative proportions compared to sporadic AD. Similarly, the APOE ε4 allele also showed decreased neuronal and increased astrocyte relative proportions compared to AD non-carriers. In contrast, carriers of variants in TREM2 risk showed a lower degree of neuronal loss compared to matched AD cases in multiple independent studies. These findings suggest that genetic risk factors associated with AD etiology have a specific effect on the cellular composition of AD brains. The digital deconvolution approach provides an enhanced understanding of the fundamental molecular mechanisms underlying neurodegeneration, enabling the analysis of large bulk RNA-sequencing studies for cell composition. It also suggests that correcting for the cellular structure when performing transcriptomic analysis will lead to novel insights of AD.

With deconvolution methods to delineate cell population changes in disease condition, it would help interpret transcriptomics results and reveal transcriptional changes in a cell type specific manner. One application demonstrated in this dissertation work is to use cell type proportion as quantitative trait to identify genetic factors associated with cellular composition changes. I performed cell type QTL analysis and identified a common pathway associated with neuronal protection underlying aging brains in the presence or absence of neurodegenerative disease symptoms. A protective variant of TMEM106B, which was previously identified with a protective effect in FTD, was identified to be associated with neuronal proportion in aging brains, suggesting a common pathway underlying neuronal protection and cognitive reservation in elderly. This extended analysis yield from deconvolution results demonstrated one promising direction of using deconvolution followed by cell type QTL analysis in identifying new genes or pathways underlying neurodegenerative or aging brains.
To understand the complexity of the brain under disease condition, network analysis as a large-scale system-level approach provides unbiased and data-driven view to identify gene-gene interactions altered by disease status. Using network analysis, I replicated and reconfirmed the co-expression pattern between MS4A gene cluster and TREM2 in sporadic AD, from which further evidence was inferred from Bayesian network analysis to show that MS4A4A might be a potential regulator of TREM2 that is validated by in-vitro experiments. In Autosomal Dominant AD (ADAD) cohort, disrupted and acquired genes were identified from PSEN1 mutation carriers. Among these genes, previously identified AD risk genes and pathways were revealed along with novel findings. These results demonstrated the great potential of applying network approach in identifying disease associated genes and the interactions among them.

To conclude the dissertation work from methodological, empirical, and theoretical levels, deconvolution pipeline for bulk RNA-Seq, cell type QTL analysis, and network analysis approaches were applied to understand transcriptome changes underlying disease etiology. From which previous AD related findings were replicated that validated the methods, and novel genes and pathways were identified as potential new therapeutic targets. Based on prior knowledge and empirical evidence observed from this dissertation work, a model is proposed to explain how genetic factors are assembled as a highly interconnected interactome network to affect proteinopathy observed in neurodegenerative disorders, that cause cellular composition changes in the brain, which ultimately leads to cognitive and functional deficits observed in AD patients.
Chapter 1: Introduction and Overview
1.1 The Alzheimer’s Disease discovery and its impact nowadays

In 1901, Alois Alzheimer, a German psychiatrist and a lecturer at the Munich University Hospital received a patient case of a 51 years female named Auguste D[129]. She was sent by her husband describing her symptoms as paranoid, progressive sleep and memory disturbance, aggressiveness, crying, and confusion. This lady was admitted to the Community Psychiatric Hospital at Frankfurt am Main, and remained impatient until her death in 1906. The brain material of her autopsy was sent to Alzheimer for examination, from which he observed distinctive plaques and neurofibrillary tangles in the histology, which were later identified as pathological hallmarks of Alzheimer’s disease. In 1906, Alzheimer presented his finding of this “peculiar” dementia case in the 37th Meeting of South-West German Psychiatrists in Tubingen. Although at the meeting it did not spur much interests from the audience, Alzheimer’s finding was included as “Alzheimer's disease” in the 3th edition of his coworker Emil Kraepelin’s text ‘Psychiatrie’ in 1910[129].

For the past one hundred years since its first diagnosis, Alzheimer’s Disease (AD) is like a shadow that never leaves, and it also grows larger as human life expectancy increases as age being its most important risk factor. In 2019, there are estimated 5.8 million Americans living with AD affecting 16 million family members. At certain stage of the disease course, patients with inability of maintaining their daily functioning highly depend on caregivers, primarily family caregivers, that incur estimated 18.4 billion unpaid hours of cares, which is equivalent to 232 billion dollars[15, 69]. These huge economic burdens and inevitable emotional distress on the family and the society would also increase as the number of AD affected population could triple by 2050[121].
As the sixth leading cause of death in the U.S., the disease course usually last from 7 to 10 years on average before the consequential death[133]. Due to progressive neuronal death in the affected brain regions, apart from cognitive functions it will also disable movement functions of the patients with the results being long term bed-bound and later having swallowing problems that ultimately lead to organ failure or lethal aspiration pneumonia[15].

1.2 Cognitive deficits as the primary clinical symptoms and related measurements

Among all the neurodegenerative disorder that would result in dementia, Alzheimer’s Disease is the most common form[23]. Impaired declarative memory is usually the first noticeable sign but sometimes it could also be other executive functions such as planning or problem-solving skills. More detailed cognitive deficit rating scales, such as the clinical dementia rating[189] (CDR), have been developed to further categorize cognitive performance for both clinical and research purposes. CDR = 0 is considered as normal without dementia; CDR = 0.5 is very mild dementia; CDR = 1 is mild dementia; CDR = 2 is moderate dementia; CDR = 3 is severe dementia.

As our knowledge about neurodegenerative disorders accumulates, neuropsychological tests also evolve to optimize their diagnostic and prognostic utilities. Efforts have been spent on ensuring the test construct validity and stability with appropriate norms[88]. Test construct validity hinges heavily on an accurate and unambiguous design that projects the clinical batteries to their designated cognitive domains to ensure they are measuring what they are designed to measure. The assessment criteria of AD developed by the Consortium to Establish a Registry for Alzheimer’s Disease (CERAD) integrates evidence from multiple modalities including clinical,
behavioral, demographical, neuropsychological, neuropathological, neuroimaging, family history, and postmortem materials to standardize and ensure accurate diagnosis[91].

1.3 Diagnostic evidence from genetic, imaging, blood and CSF biomarkers

Symptoms, such as memory loss or difficulties with executive functions, are usually what AD patients first complained to the physicians, however, pathologic changes have developed decades (10-20 years) before cognitive symptoms onset[138, 211]. To capture preclinical stages and the disease development trajectory and dynamics, a variety of biomarkers or diagnostic evidence based on genetic, CSF, blood, and imaging biomarkers have been developed to facilitate early detection and differentiation among different dementia sub forms.

1.3.1 Genetics

Amyloid β is one of the two pathological hallmarks of AD, which was first isolated from a late onset AD (LOAD) patient[105]. Later, the same authors isolated cerebrovascular amyloid protein from Down’s syndrome, a disease caused by the presence of all or part of a third copy of chromosome 21(trisomy 21). Because of the close resemblance of the two proteins and the cerebrovascular amyloid protein discovery in Down’s syndrome, the authors accurately predicted that the amyloid β gene might be located on chromosome 21[104]. Three amyloid β related genes amyloid-beta precursor protein (APP), presenilin 1 (PSEN1), presenilin 2 (PSEN2) were identified associated with familial AD with in a Mendelian dominance pattern and high penetrance[148]. Then an amyloid cascade was hypothesized (Figure 1.1) suggesting over accumulation or failed clearance of amyloid β is the central event in the pathogenesis of AD, which led to neuronal and synaptic dysfunction, and ultimately to cognitive deficits[106]. Many statements surround the amyloid centered theory have been fulfilled but one issue raised
regarding whether amyloid β being the cause or the consequence of AD. If amyloid accumulation is the leading cause, then drugs targeting clearance of amyloid should ameliorate the symptom. However, so far, none of the drug developments targeting amyloid pathway is successful, which may suggest the alternative hypothesis that amyloid accumulation might be the consequence or by-product of AD.

A second doubt surround the amyloid pathway comes from the differences in inheritance mechanism between sporadic and familial AD. Unlike familial AD which has a clear Mendelian inheritance pattern with three major gene players and an early onset of disease manifestation, sporadic AD is late onset and attributed to complex traits with multiple risk genes located throughout the genome[162]. It seems that genes with rare variants such as APP, PSEN1, PSEN2 in familial AD exert high risk effects to familial AD, whereas many genes with common variants exert medium or low risk effects to sporadic AD (Figure 1.2). For sporadic AD, apolipoprotein E (APOE) is the gene with the largest and dosage dependent effect[84]. APOE has three major

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**Figure 1.1 Amyloid cascade hypothesis.** Image reproduced from Blennow et al.[29] with permission.
alleles, protective allele ε2, common allele ε3, and risk allele ε4. In Caucasian cohorts, carriers of two ε4 alleles have increased risk of 14.9 relative to two ε3 alleles. Walking down to the risk ladder, one ε4 allele renders increased risk of 2.6 for ε2/ε4, and increased risk of 3.2 for ε3/ε4. Carriers with protective allele ε2 have reduced risk of 0.6 in both ε2/ε2 and ε2/ε3[84]. APOE influence LOAD risk in an amyloid dependent manner[235, 252]. The rare TREM2 variant p.R47H (rs75932628) carriers exhibit increased AD risk by a range from 1.7-fold to 3.4-fold[112, 212]. This gene is related to microglia and immune system through neuroinflammation[79]. The involvement of immune system leads to another hypothesis for AD surround inflammation and infection with microbial triggers, for example herpes infection[219] and oral P. gingivalis infection[75, 203]. Instead of thinking genetic variants as disease causing factors, is it possible that the genetic vulnerability that failed to protect the brain from insult is the cause of sporadic AD? For example, the vulnerability of blood brain barrier[254] and neuroinflammation triggered by microbial, stress, or even lack of sleep[68] initiate amyloid protein accumulation, then leads to neuronal death and subsequent cognitive deficits. The shift of thinking paradigm may drive therapeutic strategy and drug design switching from amyloid clearance to anti-inflammation or boost immune resilience to insults. Apart from researches focusing on protein coding genes, investigation of non-coding RNA in neurodegeneration[232] and 3D spatial structure of genome[218] may also shed light on figuring out the mechanism underlying AD.

As shown in Figure 1.2, AD has a substantial but heterogeneous genetic component. The rare mutations in the APP, PSEN1 and PSEN2[64, 238] that cause autosomal dominant AD (ADAD) only account for 1-2% of overall AD cases. There are also early-onset AD cases with unknown genetic risk factors that remain elusive. Apart from early-onset AD, the most common
manifestation of AD presents late-onset (LOAD) and accounts for the majority of the cases (90-95%). Despite appearing sporadic in nature, a complex genetic architecture underlies LOAD risk. \textit{APOE} ε4 as discussed above is the most common genetic risk factor. In addition, recent whole genome and whole exome analysis have identified rare coding variants in \textit{TREM2}[26, 113], \textit{PLD3}[58], \textit{ABCA7}[65, 249] and \textit{SORL1}[86, 226] that are associated with AD and confer risk comparable to that of carrying one \textit{APOE} ε4 allele. Besides age at onset, the clinical presentations of LOAD and ADAD are remarkably similar with an amnestic and cognitive impairment phenotype[230, 258]. A minor fraction of cases of ADAD have additional neurological findings, sometimes also seen in LOAD[230, 258]. Twin studies have estimated that the heritability of Alzheimer's disease is about 0.74 and argued that unexplained variance is due to environmental factors[101]. So far, genetic studies have identified around 30 common and rare genetic variants that contribute to the AD phenotypes; however, these genes with disease susceptibility only explain a small proportion of the genetic heritability of the AD population. The remaining unexplained heritability has been named as missing heritability[80, 177, 178]. In \textbf{Chapter 4}, an omnigenic model will be discussed from a network perspective to explain a potential cause of the missing heritability problem.
Biomarkers are objective measures of biological or pathogenic changes that can be used as diagnostic, prognostic, or disease progression measurement tools. Cerebrospinal fluid (CSF) is a clear and colorless fluid that the brain and spinal cord are immersed in. Due to its direct contact with the extracellular space of the brain, it is optimal as a biomarker to capture biological or pathogenic changes in the brain. CSF biomarkers can be divided into basic and core biomarkers. Basic biomarkers measure basic brain function that might be changed in AD condition but not specific to AD, which include measurements of blood brain barrier and immune system response to chronic inflammation (Table 1.1 basic biomarker section). The core biomarkers measure AD specific molecular pathology that is specific to AD, including APP metabolism and amyloid deposition, tau phosphorylation and axonal degeneration (Table 1.1 core biomarker section).

Figure 1.2 AD gene risk allele frequency and risk effect. Image from Karch and Goate[148] with permission.

1.3.2 CSF and plasma biomarkers

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Amyloid β proteins of different length are cleaved from APP protein by beta-secretase and gamma-secretase (Figure 1.3), and Aβ40, Aβ42, and Aβ40:42 ratios are primarily measured as biomarkers. Six different tau isoforms can be divided through alternative splicing from exon 2, 3, and 10 of MAPT gene (Figure 1.4a). There are a number of phosphorylation sites of threonine and serine across the tau isoforms but the commonly referred as phosphorylated tau levels are measured from Thr181 or Thr231 phosphorylation sites (Figure 1.4b). Other CSF biomarkers related to neuronal and synaptic proteins, for example, visinin-like protein 1 (VLP-1) and synaptosomal-associated protein 25 (SNAP-25), and oxidative stress markers such as F2-isoprostanes were also be able to differentiate AD from controls.

![Figure 1.3 APP protein cleavage and amyloid β proteins.](image)

*Image from Blennow et al.[29] with permission.*

<table>
<thead>
<tr>
<th>Table 1.1 CSF biomarkers</th>
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<tr>
<td><strong>Biomarker</strong></td>
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</tr>
<tr>
<td>Basic biomarker</td>
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<tr>
<td>CSF cell count</td>
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<tr>
<td>CSF: serum albumin ratio</td>
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<td>IgG or IgM index or oligoclonal bands</td>
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<tr>
<td>Aβ 1-42</td>
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<tr>
<td>p-tau181 and p-tau231</td>
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<td>total tau</td>
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CSF biomarkers have shown great potential in measuring pre-symptomatic changes before the plaque becomes too widespread or the proceeding of irreversible neurodegeneration. However, collection of CSF through lumbar puncture is invasive with potential risks of post-lumbar puncture headache, back discomfort or pain, bleeding, and brainstem herniation. Therefore, developments of non-invasive biomarkers, such as plasma derived biomarker, have been pursuit to look for alternative sources other than CSF, for example plasma Aβ\textsubscript{1-42}[180], Aβ\textsubscript{40} to Aβ\textsubscript{42} ratio[109], APP\textsubscript{669-711} to Aβ\textsubscript{42} ratio[195], α 2-Macroglobulin (α2M) and complement factor H (CFH)[135], neurofilament light protein (NFL)[179].

1.3.3 Imaging

As amyloid β aggregation being directly related to AD and potentially a predictor of AD decades before cognitive deficits, Aβ PET imaging has been used clinically as a diagnosis tool and prognosis measurement[270]. Although the Aβ’s role in AD is still under debate, it has been shown that amyloid β deposition proceeds neuronal loss or cerebral atrophy observed on MRI. Five different amyloid β tracers showed that the frontal, temporal and posterior cingulate cortices
showed the highest retention rate for Aβ (Figure 1.5) that correlates with regional amyloid β plaque density and the sequence of amyloid β deposition found in post-mortem brains in sporadic AD[32]. Noticeably, the pattern of amyloid β tracer retention in familial AD mainly located in striatal region which is different from the patterns of sporadic AD. Based on the retention patterns observed, amyloid β imaging could also be used to differentiate sporadic AD from dementia with Lewy bodies (DLB) and early-onset AD from frontotemporal lobar degeneration (FTLD), because DLB exhibits a posterior retention pattern that sporadic AD does not have, and FTLD should not have C-PIB retention. However, it is worth noticing that about 25% cognitive normal elderly also have fibrillary Aβ deposition[185], which had been observed long before the amyloid β PET imaging era[216].

Figure 1.5 Amyloid β PET radiotracer imaging. Surface projection images obtained from five AD patients with different amyloid beta PET radiotracer. Five images showed consistent pattern with highest retention in frontal, temporal and posterior cingulate cortices representing amyloid β deposition in the brain. Image from Villemagne et al.[270] with permission.
Similar to amyloid β imaging, tau imaging also showed consistent tau tracer retention pattern (Figure 1.6) with post-mortem studies, besides it is more correlated with neuronal injury markers. As opposed to amyloid β imaging which focusing on total amyloid β load, regional tau distribution in terms of density and topological distribution of tau provide more information in disease progression than total tau load.

Apart from tracers based PET that requires tracer injection into the patients, gradient recalled echo MRI based approach post-processing method, Gradient Echo Contrast Imaging (GEPCI), has been developed utilizing transverse relaxation rate constant to avoid tracer injection. The GEPCI metrics showed strong correlation with both amyloid β accumulation measured from PET and cognitive performance[292]. Another non-tracer imaging technique based on functional connectivity MRI also showed success in differentiating APOE4+ from APOE4- carriers in the absence of amyloid deposition[240] suggesting early genetic effect can be measured with functional connectivity MRI. Strong evidences suggested the default mode network is strongly associated with AD[17, 111, 116, 241].

Since 2011, the diagnosis guideline for AD in the U.S. had been revised from the 1984 diagnostic criteria, which was mainly based on the clinical judgement of the patient’s symptoms, to incorporate biomarker tests[15]. An A/T/N system have been proposed to integrate multiple biomarker modalities: “A” refers to β-amyloid related biomarker including amyloid PET or CSF Aβ42; “T” refers to tau related biomarker including CSF phosphor tau or tau PET; and “N” refers to non-specific neurodegeneration or neuronal injury biomarkers including 18F-FDG PET, structural MRI or CSF total tau[137]. With either positive or negative binary traits defined by respective cutoffs within each category, a biomarker profile integrating multimodal measurements can be established for the subject to inform diagnosis[136]. As mentioned above,
tau imaging result is highly correlated with neuronal function biomarker $^{18}$F-FDG PET, thus one potential problem with ATN is that the integrative approach includes highly correlated metrics may incur a cost of redundant tests or repetitive information. Besides, the binary traits may over simplify the complexity of AD manifestation as opposed to a more quantitative approach.

![Figure 1.6 Tau radiotracer imaging](image)

**Figure 1.6 Tau radiotracer imaging.** Representative PET images with three different tau radiotracers. Top row is sagittal view; center row is transverse view; bottom row is coronal view. Compared to healthy elderly controls (HC), AD patients showed tracer retention in mesial temporal, temporoparietal and posterior cingulate cortical regions. $^{18}$F-THK5351-PET and $^{18}$F-flortaucipir-PET in HC show ‘off-target’ retention in the striatum. $^{18}$F-THK5351-PET also show tracer retention in the striatum. Image from Villemagne et al.[270] with permission.

### 1.4 Neuropathological verification for postmortem assessment of AD

Mostly for research purposes, neuropathological assessments are performed during autopsy on post mortem materials to verify the clinical diagnosis of AD. The CERAD neuropathology criteria contains gross and microscopic findings focusing on hippocampus, amygdala and various cortical regions[91, 186]. They use a semi-quantitative approach to assess frequency of senile plaques, including both neuritic plaques relative to the patient’s age and diffuse plaques, neurofibrillary tangles, and others such as cerebrovascular changes. From those
together with clinical history, a categorical assessment result will be derived to report the

certainty of AD diagnosis, and they are definite, probable, possible, or no evidence of AD.

Other commonly used neuropathological assessment with slightly different focuses are
Braak and Braak[34], Khachaturian[150], NIA-Reagan Institute[5], and the Tierney A3[259]
criteria. Braak and Braak staging focuses on the distribution patterns of neurofibrillary tangles
and neuropil threads[34], which is divided into six stages – stage I and II are characterized by
either mild or severe alteration of the transentorhinal region; stage III and IV are marked by
conspicuous changes in both transentorhinal region and entorhinal cortex; stage V and VI
include destruction of all isocortical associated regions. Khachaturian[150] documented
consensus criteria of AD diagnosis reached upon by the neuropathology panel at the “research
workshop on the diagnosis of Alzheimer’s Disease” organized by National Institute of Aging
(NIA) in 1983. These criteria focus on microscopic findings in frontal, temporal, and parietal
lobes, the amygdala, the hippocampal formation, the basal ganglia, the substantia nigra, the
cerebellar cortex, and the spinal cord. The number of amyloid plaque and neurofibrillary tangles
per field for different age ranges are specified for AD diagnosis. Another more recent consensus
recommendation of postmortem diagnostic criteria for AD are proposed to reassesses the
previous criteria documented in Khachaturian[150]. This meeting was led by both NIA and
emphasize on the heterogeneous clinicopathological characteristics of AD, thus the diagnosis are
only probabilistic rather than deterministic statements in any given patient based on both
CERAD and Braak and Braak staging criteria. Besides, there might be other pathological process
involved along with AD that contribute to dementia, for example stroke, Parkinson’s disease,
progressive supranuclear palsy, and etc. A study that compared different pathological criteria
found the CERAD category of definite AD closely resemble the cases that fulfil the Tierney A3[259] AD criteria[194].

1.5 Relation to other neurodegenerative disorders

Under the umbrella term of neurodegenerative disease resulted from neuronal loss, patients suffering from various cognitive or motor deficits are categorized into different arbitrarily defined diseases based on their clinical manifestations. Despite distinct symptoms and affected brain regions (Table 1.2), different neurodegenerative diseases share some common features that may suggest potential shared mechanisms underlying disease etiology[99]. For example, the two major clinical manifestations, cognitive deficit and motor deficit, divide the realm into two halves. Age is the most important risk factor for all of the neurodegenerative diseases. Aggregation and progression of misfolded proteins are also involved in all of them, although being the cause or the result of the disease is still under debate. The common features suggest common pathways being altered in neurodegenerative diseases, including protein quality control, the autophagy-lysosome pathway, mitochondria homeostasis, protein seeding and propagation of stress granules, and synaptic toxicity and network dysfunction[99]. Genetically, MAPT gene (microtubule associated protein tau) only plays a modest role in sporadic AD but a substantial role in Frontotemporal Dementia (FTD) and Progressive Supranuclear Palsy (PSP). The most important susceptibility region for late-onset AD surrounding the APOE gene is involved in non-AD neurodegenerative disorders and conditions[62], such as Lewy body dementias(LBD)[25, 284], Parkinson's disease (PD)[40], amyloid angiopathy[25, 110, 285], TDP-43 proteinopathy[283], hippocampal sclerosis[25, 83, 283].
1.6 Dissertation Overview

As discussed above, unlike the rare familial Mendelian dominant AD, late onset AD inherited as complex traits are more common in the population and resulted from dozens of variants involving genes distributed across the whole genome. To tackle the complex interaction in AD, the primary purpose of this dissertation is to apply integrative analysis approaches to demystify and obtain a more accurate and comprehensible picture of AD etiology.

Alzheimer’s disease is characterized by neuronal loss and astrocytosis in the cerebral cortex. However, the specific effects that pathological mutations and coding variants associated with AD have on the cellular composition of the brain are often ignored. In chapter 2, to investigate cerebral cortex cell-type population structure I developed an in-silico deconvolution method to infer cellular composition from RNA-Seq. I firstly assembled a reference panel to model the transcriptomic signature of neurons, astrocytes, oligodendrocytes and microglia. The panel was created by analyzing expression data from purified cell lines. I evaluated alternative digital deconvolution methods and selected the best performing ones for my primary analyses. I tested the digital deconvolution accuracy on induced pluripotent stem cell (iPSC) derived neurons/microglia cells and neuronal Translating Ribosome Affinity Purification followed by RNA-Seq. Finally, I verified its accuracy by creating artificial admixture with pre-defined
cellular proportions. Once the deconvolution approach was optimized, I calculated the cell proportion in AD cases and controls from the different brain regions of LOAD and ADAD datasets. I found that neuronal and astrocyte relative proportions differ between healthy and diseased brains and also among AD cases that carry specific genetic risk variants. Brain carriers of pathogenic mutations in \textit{APP}, \textit{PSEN1} or \textit{PSEN2} presented lower neurons and higher astrocytes relative proportions compared to sporadic AD. Similarly, the \textit{APOE ε4} allele also showed decreased neuronal and increased astrocyte relative proportions compared to AD non-carriers. On the contrary, carriers of variants in \textit{TREM2} risk showed a lower degree of neuronal loss than matched AD cases in multiple independent studies. These findings suggest that genetic risk factors associated with AD etiology have a specific imprinting in the cellular composition of AD brains.

In \textbf{chapter 3}, I utilized cell-type proportions inferred from deconvolution procedure as disease status proxy to identify new genetic variants related to AD. Instead of using disease phenotype, studies which used endo-phenotypes, such as CSF APOE levels[59], CSF amyloid-β, tau, and phosphorylated tau (ptau\textsubscript{181})[71], and AD proxy[174] have successfully uncovered other variants associated with AD. Using cell type composition inferred from RNA-Seq data as a disease status proxy, I performed cell type association analysis to identify potential new locus that are related to cellular population changes in disease cohort. We imputed and merged genotyping data from seven studies (five centered on neurodegeneration; two focused on schizophrenia and multiple tissue controls), and derived major CNS cell type proportions as described in chapter 2 from cortical RNA-Seq data. Neuronal proportion were normalized by subtracting the mean from each tissue deconvolution results after removing outliers. I identified a variant rs1990621 located in the \textit{TMEM106B} gene region significantly associated with neuronal
proportion in cortical RNA-Seq dataset. This variant is in high LD with rs1990622 \( (r^2 = 0.98) \) which was previously identified as a protective variant in FTD cohorts\[266\]. In conclusion, I have identified a variant associated with neuronal proportion with potential protective effect in neurodegeneration disorders.

In Chapter 4, using network analysis I replicated and reconfirmed the co-expression pattern between \textit{MS4A} gene cluster and \textit{TREM2} in sporadic AD, from which further evidence was inferred from Bayesian network analysis to show that \textit{MS4A4A} might be a potential regulator of \textit{TREM2} that is validated by \textit{in-vitro} experiments. In Autosomal Dominant AD (ADAD) cohort, disrupted and acquired genes were identified from \textit{PSEN1} mutation carriers. Among the genes, previous identified AD related gene and pathways were revealed together with novel findings. These results demonstrated the great potential of applying network approach in identifying disease associated genes and the interactions among them.

In conclusion, contribution from this dissertation work to AD research is summarized in Chapter 5, and future directions in research to facilitate diagnosis, intervention, and disease-modifying therapies are also discussed in the context of this dissertation work.
Chapter 2: Genetic variants associated with Alzheimer’s disease confer different cerebral cortex cell-type population structure

This chapter was adapted from:

2.1 Abstract

**Background:** Alzheimer's disease (AD) is characterized by neuronal loss and astrocytosis in the cerebral cortex. However, the specific effects that pathological mutations and coding variants associated with AD have on the cellular composition of the brain are often ignored.

**Methods:** I developed and optimized a cell-type-specific expression reference panel and employed digital deconvolution methods to determine brain cellular distribution in three independent transcriptomic studies.

**Results:** I found that neuronal and astrocyte relative proportions differ between healthy and diseased brains and also among AD cases that carry specific genetic risk variants. Brain carriers of pathogenic mutations in *APP*, *PSEN1*, or *PSEN2* presented lower neuron and higher astrocyte relative proportions compared to sporadic AD. Similarly, the *APOE ε4* allele also showed decreased neuronal and increased astrocyte relative proportions compared to AD non-carriers. In contrast, carriers of variants in *TREM2* risk showed a lower degree of neuronal loss compared to matched AD cases in multiple independent studies.

**Conclusions:** These findings suggest that genetic risk factors associated with AD etiology have a specific imprinting in the cellular composition of AD brains. My digital deconvolution method provides an enhanced understanding of the fundamental molecular mechanisms underlying neurodegeneration, enabling the analysis of large bulk RNA-sequencing studies for cell composition and suggests that correcting for the cellular structure when performing transcriptomic analysis will lead to novel insights of AD.
2.2 Introduction

2.2.1 Altered cellular composition confounds downstream transcriptomic analysis

Altered cellular composition is associated with AD progression and decline in cognition. Neuronal loss in the hippocampus is characteristic in the initial stages of AD, which could explain early memory disturbances[205, 282]. As the disease progresses, neuronal death is observed throughout the cerebral cortex. Furthermore, ~25% of individuals who die by ~75 years of age who were cognitively normal also presented substantial cerebral lesions that resemble AD pathology, including amyloid plaque, NFTs, and neuronal loss[132]. Thus, the identification of the brain cellular population structure is essential for understanding neurodegenerative disease progression[107]. However, stereology protocols for counting neurons can be tedious, require extensive training and are susceptible to technical artifacts which may lead to biased quantification of cell-type distributions[107].

Recently there has been a growing interest in understanding the transcriptomic changes attributed to AD[9, 46, 98, 184, 197, 206, 247, 287], as these may point to underlying molecular mechanisms of disease. These studies are typically designed to analyze the expression profiles of large cohorts ascertained from homogenized regions of the brain (e.g. bulk RNA-Seq) of affected and control donors. However, bulk RNA-Seq captures the gene expression of all of the constituent cells in the sampled tissue, and the altered cellular composition associated with AD has been reported to confound downstream analyses[247].

2.2.2 Digital deconvolution approach

Digital deconvolution approaches enhance the interrogation of expression profiles to identify the cellular population structure of individual samples, alleviating the requirement of
additional neurostereology procedures. These approaches have been developed, tested and applied to ascertain cellular composition altered in many traits[157, 199, 242, 293]. However, digital deconvolution has not been applied to identify the cellular population structure from RNA-Seq from human brain of AD cases and controls. Technical constraints restrict the dissociation of cells from the brains for very specific conditions[38, 290, 291]. Nevertheless, a limited number of RNA-Seq from isolated cell populations from the brain have been generated[38, 290, 291]. Using these resources, I am now able to generate a reference panel for digital deconvolution of human brain bulk RNA-Seq data.

I sought to investigate the cellular population structure in AD by analyzing RNA-Seq from multiple brain regions of LOAD participants. To do so, I assembled a novel brain reference panel and evaluated the accuracy of digital deconvolution methods by analyzing additional cell-type specific RNA-Seq samples and by creating synthetic admixtures with defined cellular distributions. Then I analyzed large cohorts of pathologically confirmed AD cases and controls (N = 613) and verified that it predicts cellular distribution patterns consistent with neurodegeneration. Finally, I generated RNA-Seq from the parietal lobe of participants from the Charles F. and Joanne Knight Alzheimer's Disease Research Center (Knight-ADRC)[153], including non-demented controls, LOAD cases, with enriched proportions of carriers of high-risk coding variants associated with AD, and also ADAD from The Dominantly Inherited Alzheimer Network[72] (DIAN). I compared the cell composition in ADAD and LOAD; and also evaluated differences among carriers of coding high-risk variants in PLD3, TREM2 and APOE ε4. My findings indicated that cell-type composition differs among carriers of specific genetic risk factors, which might be revealing distinct pathogenic mechanisms contributing to disease etiology.
2.3 Methods

2.3.1 Subjects and Samples

DIAN and Knight-ADRC

Parietal lobe tissue of post-mortem brain was obtained with informed consent for research use and were approved by Washington University in St. Louis review board. RNA was extracted from frozen brain using Tissue Lyser LT and RNeasy Mini Kit (Qiagen, Hilden, Germany). RNA-Seq Paired end reads with read length of 2×150bp were generated using Illumina HiSeq 4000 with a mean coverage of 80 million reads per sample (Table 2.1; Table 2.2). RNA-Seq was generated for 19 brains from The Dominantly Inherited Alzheimer Network (DIAN), 84 brains with late-onset AD and 16 non-demented controls from The Charles F. and Joanne Knight Alzheimer's Disease Research Center (Knight ADRC)[153]. The AD brains selected from the Knight-ADRC are enriched for carrier of variants in TREM2 (N=20; Table 2.2) and PLD3 (N=33; Table 2.2). The clinical status of participants was neuropathologically confirmed[187]. We identified three additional participants from the Knight-ADRC study with PSEN1 (p.A79V, p.I143T and p.S170F) mutations. CDR scores were obtained during regular visits throughout the study prior to the subject’s decease[190]. A range of other pathological measurement were collected during autopsy including Braak staging, as previously described[35].
RNA was extracted from frozen brain tissues using Tissue Lyser LT and RNeasy Mini Kit (Qiagen, Hilden, Germany) following the manufacturer’s instruction. RIN (RNA integrity) and DV200 were measured with RNA 6000 Pico Assay using Bioanalyzer 2100 (Agilent Technologies). The RIN is determined by the software on the Bioanalyzer taking into account the entire electrophoretic trace of the RNA including the presence or absence of degradation products. The DV200 value is defined as the percentage of nucleotides greater than 200nt. RIN and DV200 for all the samples can be found on Table 2.2. The yield of each sample is determined by the Quant-iT RNA Assay (Life Technologies) on the Qubit Fluorometer (Fisher Scientific). The cDNA library was prepared with the TruSeq Stranded Total RNA Sample Prep with Ribo-Zero Gold kit (Illumina) and then sequenced by HiSeq 4000 (Illumina) using 2×150 paired end reads at McDonnell Genome Institute, Washington University in St. Louis with a

| Table 2.1 Demographics and disease status of cohorts from four brain bank resources. |
|---------------------------------|-----------------|--------------|-------------|----------------|
| Sample Size                     | Mayo<sup>a</sup> | MSBB<sup>b</sup> | DIAN        | Knight-ADRC   |
| Age                             | 83 ± 7.77       | 83.3 ± 7.55  | 50.6 ± 7.06 | 85.1 ± 9.78   |
| % Male                          | 45.5            | 36           | 68.4        | 38.8          |
| % <i>APOE</i> e4+                | 33.2            | 31.7         | 14.3        | 45.6          |
| Brain weight                    | -               | -            | 1187.7 ± 184.5 | 1138.1 ± 142.5 |
| AD<sup>c</sup>                  | 82              | 135          | 19          | 87            |
| PA<sup>d</sup>                  | 29              | 0            | 0           | 0             |
| Control                         | 80              | 85           | 0           | 0             |
| CDR = 0                         | -               | 40           | 0           | 13            |
| CDR = 0.5                       | -               | 40           | 0           | 9             |
| CDR = 1                         | -               | 30           | 2           | 11            |
| CDR = 2                         | -               | 44           | 4           | 14            |
| CDR = 3                         | -               | 146          | 1           | 56            |

<sup>a</sup> Mayo stands for Mayo Clinic.

<sup>b</sup> MSBB stands for Mount Sinai Brain Bank.

<sup>c</sup> AD stands for Alzheimer’s Disease.

<sup>d</sup> PA stands for pathological aging (amyloid plaques but no tau tangles).

<sup>e</sup> CDR stands for clinical dementia rating for available samples.
mean of 58.14 ± 8.62 million reads. Number of reads and other QC metrics can be found in Table 2.2.

Mayo Clinic Brain Bank

Mayo Clinic Brain Bank RNA-Seq was accessed from the AMP-AD portal (synapse ID = 5550404; accessed January 2017) (Table 2.1). Paired end reads of 2×101bp were generated by Illumina HiSeq 2000 sequencers for an average of 134.9 million reads per sample. Neuropathology criteria, quality control procedures, RNA extraction and sequencing details are explained elsewhere[9].
Table 2.2 Demographics and AD mutation carriers of DIAN and Knight-ADRC cohorts.

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<td>RIN</td>
<td>5.69 ± 1.13</td>
<td>6.44 ± 1.16</td>
<td>6.71 ± 1.18</td>
<td>6.34 ± 1.19</td>
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<td>DV200</td>
<td>86.59 ± 4.12</td>
<td>89.48 ± 3.85</td>
<td>91.19 ± 2.54</td>
<td>89.18 ± 3.97</td>
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<td>PMI</td>
<td>14.57 ± 10.29</td>
<td>13.05 ± 6.66</td>
<td>10.52 ± 6.09</td>
<td>12.99 ± 7.4</td>
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<td>Age</td>
<td>51.27 ± 11.13</td>
<td>85.72 ± 6.83</td>
<td>87.08 ± 10.2</td>
<td>79.69 ± 15.67</td>
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<td>Male %</td>
<td>0.64</td>
<td>0.39</td>
<td>0.38</td>
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<tr>
<td>APOE4+ %</td>
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<td>0.52</td>
<td>0.06</td>
<td>0.44</td>
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<td>CDR</td>
<td>2.2 ± 0.79</td>
<td>2.37 ± 0.93</td>
<td>0.22 ± 0.31</td>
<td>2.04 ± 1.14</td>
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<td>Braak</td>
<td>5.94 ± 0.24</td>
<td>4.84 ± 1.29</td>
<td>1.93 ± 0.88</td>
<td>4.61 ± 1.62</td>
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<td>Number of Total Reads (Million)</td>
<td>60.92 ± 5.6</td>
<td>57.7 ± 9.28</td>
<td>56.6 ± 7.98</td>
<td>58.14 ± 8.62</td>
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<tr>
<td>Uniquely Mapped Reads %</td>
<td>79.72 ± 4.28</td>
<td>80.74 ± 4.49</td>
<td>81.06 ± 5.96</td>
<td>80.6 ± 4.65</td>
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<td>Mapped to Multiple Loci Reads %</td>
<td>16.39 ± 2.1</td>
<td>15.56 ± 2.2</td>
<td>15.08 ± 3.3</td>
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<td>Disease Status</td>
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<td>Sporadic AD</td>
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*There are two Knight-ADRC subjects that carry both PLD3 and UNC5C variants.
RNA-Seq based transcriptome data was generated from post-mortem brain tissue collected from cerebellum (189 samples) and temporal cortex (191 samples) of Caucasian subjects[2, 9]. RNA was extracted using Trizol® reagent and cleaned with Qiagen RNeasy. RIN measurement was performed with Agilent Technologies 2100 Bioanalyzer. Samples with RIN greater than 5 were included. Library was prepared by Mayo Clinic Medical Genome Facility Gene Expression and Sequencing Cores with TruSeq RNA Sample Prep Kit (Illumina).

Mount Sinai Brain Bank

Mount Sinai Brain Bank RNA-Seq study was downloaded from the AMP-AD portal (synapse ID = 3157743; accessed January 2017) (Table 2.1). Single end reads of 100 nucleotides was generated by Illumina HiSeq 2500 System (Illumina, San Diego, CA) for an average of 38.7 million reads per sample[3].

This dataset contains 1030 samples collected from four post-mortem brain regions of 300 subjects: anterior prefrontal cortex (BA10), superior temporal gyrus (BA22), parahippocampal gyrus (BA36), and inferior frontal gyrus (BA44). RNA-Seq was generated using the TruSeq RNA Sample Preparation Kit v2 and Ribo-Zero rRNA removal kit (Illumina, San Diego, CA)[3].

iPSC-derived neurons

Dermal fibroblasts were obtained from skin biopsies from research participants in the Knight-ADRC (Fibroblast lines: F11362, F12455, and F13504). Human fibroblasts were reprogrammed into iPSC using non-integrating Sendai virus carrying OCT3/4, SOX2, KLF4, and cMYC[255, 265]. iPSCs were manually selected and expanded on Matrigel in mTesR1 (StemCell Technologies). iPSCs were characterized for expression of pluripotency markers by immunocytochemistry and quantitative PCR (qPCR). qPCR with probes specific to Sendai virus
were used to confirm the absence of virus in the isolated clones. All cell lines were confirmed to have a normal karyotype based on G-band karyotyping. To generate cortical neurons, iPSCs were plated in a v-bottom plate in neural induction media (StemCell Technologies; 65,000 per well) to form highly uniform neural aggregates. After 5 days, neural aggregates were transferred onto PLO/laminin-coated tissue culture plates. Neural rosettes formed over 5-7 days. The resulting neural rosettes were then isolated by enzymatic selection (StemCell Technologies) and cultured as neural progenitor cells (NPCs). NPCs were then differentiated by culturing in neural maturation medium (neurobasal medium supplemented with B27, GDNF, BDNF, cAMP). RNA was collected from the cells and sequenced following the same protocol and processing pipeline as the DIAN and Knight-ADRC dataset.

In addition, I accessed RNA-Seq data generated for iPSC-derived neurons from the Broad iPSC study[7] (Synapse ID: syn3607401). Forebrain neurons from wild-type background were generated using an embryoid body-based protocol to produce neural progenitor cells (day 17) and mature neurons (day 57 and day 100). RNA was purified using a PureLink RNA mini-kit (Life Technologies). RNA-Seq libraries were prepared using Illumina Strand Specific TruSeq protocol, and sequenced to obtain an average of 75M reads in pairs reads per sample.

**TRAP-seq mice**

All animal procedures were performed in accordance with the guidelines of Washington University's Institutional Animal Care and Use Committee. The Rosa26^{fsTRAP} mice (Gt(ROSA)26Sor^{tm1(CAG-EGFP/Rpl10a,-birA)Wtp})[294] (The Jackson Laboratory) were crossed with PV^{Cre} mice (Pvalb^{tm1(cre)Arbr})[128] (The Jackson Laboratory) to produce PV-TRAP mice directing expression of EGFP-L10a ribosomal fusion protein in parvalbumin (PV) expressing cells.
Purification of cell-type specific mRNA by translating ribosome affinity purification (TRAP) was described previously[122] with modifications. Briefly, PV-TRAP mouse brain was removed and quickly washed in ice-cold dissection buffer (1× HBSS, 2.5 mM HEPES-KOH (pH 7.3), 35 mM glucose, and 4 mM NaHCO\textsubscript{3} in RNase-free water). Barrel cortex was rapidly dissected and flash-frozen in liquid nitrogen, and then stored at -80 °C until use. Affinity matrix was prepared with 150 μl of Streptavidin MyOne T1 Dynabeads, 60 μg of Biotinylated Protein L, and 25 μg of each of GFP antibodies 19C8 and 19F7. The tissue was homogenized on ice in 1 ml of tissue-lysis buffer (20 mM HEPES KOH (pH 7.4), 150 mM KCl, 10 mM MgCl\textsubscript{2}, EDTA-free protease inhibitors, 0.5 mM DTT, 100 μg/ml cycloheximide, and 10 μl/ml rRNasin and Superasain). Homogenates were centrifuged for 10 min at 2,000 × g, 4 °C, and 1/9 sample volume of 10% NP-40 and 300 mM DHPC were added to the supernatant at final concentration of 1% (vol/vol). After incubation on ice for 5 min, the lysate was centrifuged for 10 min at 20,000 × g to pellet insolubilized material. Then 200 μl of freshly resuspended affinity matrix was added to the supernatant and incubated at 4 °C for 16–18 hours with gentle end-over-end mixing in a tube rotator. After incubation, the beads were collected with a magnet and resuspended in 1000 μl of high-salt buffer (20 mM HEPES KOH (pH 7.3), 350 mM KCl, 10 mM MgCl\textsubscript{2}, 1% NP-40, 0.5 mM DTT and 100 μg/ml cycloheximide), and collected with magnet as above. After 4 times of washing with high-salt buffer, RNA was extracted using Absolutely RNA Nanoprep Kit (Agilent Technologies) following manufacturer’s instruction. RNA quantification was measured using Qubit RNA HS Assay Kit (Life Technologies) and the integrity was determined by Bioanalyzer 2100 using an RNA Pico chip (Agilent Technologies). The cDNA library was prepared with Clontech SMARTer and then sequenced by HiSeq3000. Single end reads of 50 base pairs were generated for an average of 29.2 million reads per sample (24 samples).
iPSC-derived microglia

The data was accessed from the AMP-AD portal (Synapse ID: syn7203233). This dataset is comprised of iPSC-derived microglia (N = 10) from human primitive streak-like cells[77]. Within 30 days of differentiation, myeloid progenitors coexpressing CD14 and CX3CR1 were generated. These iPSC-derived microglia were able to perform phagocytosis and elicit ADP-induced intracellular Ca\textsuperscript{2+} transients that asserted their microglia identity as opposed to macrophage. Single-ended RNA-Seq data was generated with the Illumina HiSeq 2500 platform following the Illumina protocol.

2.3.2 RNA-Seq QC and Alignment

FastQC was applied to DIAN and Knight-ADRC RNA-Seq data to perform quality check on various aspects of sequencing quality[231]. Each category of FastQC will be explained with pass or fail examples together with summary results ascertained from the DIAN and Knight-ADRC combined dataset. QC result explanations were obtained from the developer’s website[11].

The DIAN and Knight-ADRC dataset was aligned to human GRCh37 primary assembly using Star (ver 2.5.2b)[74]. I used the primary assembly and aligned reads to the assembled chromosomes, un-localized and unplaced scaffolds, and discarded alternative haploid sequences. Sequencing metrics, including coverage, distribution of reads in the genome[4], ribosomal and mitochondrial contents and alignment quality, were further obtained by applying Picard CollectRnaSeqMetrics (ver 2.8.2) to detect sample deviation. QC results from FastQC, Star, Picard, and Salmon are merged with multi-QC software to generated integrated summary reports (Table 2.3).
Problematic samples summary:

- RIN < 5 & DV200 < 75
  - H_VY-82018_S1512310_II.H.40
  - H_VY-83774_S1512313_II.G.39
- Low Yield
  - H_VY-60410_S1511525_I.D.19
  - H_VY-62240_S1511620_IV
- High Ribosomal RNA
  - H_VY-83774_S1512313_II.G.39
  - H_VY-82018_S1512310_II.H.40
  - H_VY-9TPSKM_S1512275_I.B.15
- High Median 5’ to 3’ bias
  - H_VY-60007_S1511546_I.D.21
  - H_VY-83774_S1512313_II.G.39
- Ethnicity non-Europeans
  - H_VY-83665_S1511508_I.E.31
  - H_VY-61256_S1511537_I.E.31
  - H_VY-62275_S1511542_I.D.19
  - H_VY-11964_S1512298_I.E.29
  - H_VY-23178_S1512475_I.D.19
  - H_VY-62464_S1512484_VI.N
- Transcriptome-wise outliers
  - H_VY-F1R54Y_D1202616_I.C.18
  - H_VY-1XYTL9_D1202619_I.B.10
  - H_VY-61377_S1511495_I.D.19
  - H_VY-61245_S1512302_IV
  - H_VY-61684_S1512304_IV
  - H_VY-60007_S1511546_I.D.21
  - H_VY-83774_S1512313_II.G.39
Table 2.3 Summarized quality check results integrated with Multi-QC

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<th>Sample Name</th>
<th>Picard % RNA</th>
<th>Picard % mRNA</th>
<th>Salmon % Aligned</th>
<th>Salmon M Aligned</th>
<th>STAR % Uniquely Aligned</th>
<th>STAR M Uniquely Aligned</th>
<th>FastQC % Dups Reads</th>
<th>FastQC % GC</th>
<th>FastQC Total M Seqs</th>
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<tr>
<td>H_VY-60071_51512411_L.E.34</td>
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<tr>
<td>H_VY-60826_51511514_L.D.19</td>
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</tbody>
</table>

32
| H_VY-61648_D1202287_L1.E.32 | 0.80% | 39.00% | 41.20% | 25 | 85.70% | 42.1% | 21.4 | 83.20% | 45.9% | 52.1 | 38.70% | 45% | 121.6 |
| H_VY-61684_S1512304_IV | 0.80% | 33.40% | 36.20% | 49 | 42.00% | 45% | 117.8 |
| H_VY-61741_S1512305_IV | 0.80% | 32.40% | 35.10% | 53.3 | 39.80% | 45% | 133 |
| H_VY-64269_S1512482_L1 | 0.80% | 32.20% | 35.00% | 31.4 | 30.10% | 46% | 72.7 |
| H_VY-72221_S1511778_VLN | 0.80% | 33.20% | 36.50% | 51.5 | 35.90% | 45% | 121.9 |
| H_VY-85194_S1511391_L8.9 | 0.80% | 36.40% | 38.90% | 50.9 | 39.20% | 46% | 121.8 |
| H_VY-60230_S1511547_L0.21 | 0.70% | 34.10% | 36.00% | 51.5 | 36.60% | 45% | 124.1 |
| Pooled_RNA_2096115730 | 0.70% | 48.10% | 48.50% | 57.3 | 52.90% | 47% | 132.4 |
| H_VY-60510_S1512480_L1.E.35 | 0.60% | 34.40% | 36.40% | 53 | 40.70% | 44% | 125 |
| H_VY-60180_S1511776_VLN | 0.60% | 45.50% | 47.00% | 54.4 | 46.20% | 45% | 129.1 |
| H_VY-60974_S1512301_VLN | 0.60% | 35.50% | 37.70% | 48.3 | 34.70% | 45% | 115.7 |
| H_VY-61609_S1512483_VLN | 0.60% | 36.40% | 38.80% | 53.9 | 37.00% | 34% | 123.2 |
| H_VY-61744_S1511550_L1.E.35 | 0.50% | 34.30% | 36.60% | 43.7 | 34.90% | 45% | 101.1 |
| H_VY-62658_S1512478_VLN | 0.50% | 36.30% | 38.70% | 50.7 | 33.40% | 44% | 117.6 |
| H_VY-F11362.1d1B6_Neuron_3_S1512502_VII.0.45 | 0.50% | 41.70% | 50.00% | 37.3 | 46.70% | 47% | 90.4 |
| H_VY-F11362.1d1F10_Neuron_2_S1512504_VII.0.47 | 0.50% | 45.10% | 52.20% | 31.5 | 48.30% | 49% | 79.5 |
| H_VY-F12455.8_Astro_2_S1512507_VII.0.43 | 0.50% | 66.60% | 66.60% | 51.1 | 56.60% | 48% | 116.8 |
| H_VY-60524_S1511987_VLN | 0.40% | 36.30% | 38.50% | 53.5 | 36.60% | 45% | 127.7 |
| H_VY-61280_S1511982_V | 0.40% | 36.40% | 39.20% | 55.6 | 46.90% | 45% | 128.2 |
| H_VY-F11362.1d1B6_Neuron_1_S1512500_VII.0.45 | 0.40% | 53.40% | 54.70% | 44.4 | 52.60% | 47% | 105.4 |
| H_VY-61587_S1512303_IV | 0.30% | 33.60% | 36.80% | 55.2 | 36.20% | 44% | 124.8 |
| H_VY-F12455.8_Astro_IGF_3_S1512511_VII.0.43 | 0.30% | 67.10% | 68.60% | 50.9 | 38.00% | 43% | 115.6 |
| Pooled_RNA_2096115783 | 0.30% | 53.30% | 47.70% | 53.8 | 46.00% | 45% | 122.5 |
| H_VY-F12455.8_Astro_IGF_1_S1512509_VII.0.43 | 0.20% | 66.30% | 68.10% | 39 | 47.20% | 47% | 88.8 |
| **Average** | 2.96% | 34.38% | 36.85% | 21.4 | 80.57% | 46.8 | 41.26% | 47% | 116.2 |
Per base sequence quality

**Figure 2.1A** and **Figure 2.1B** are passed and failed example for the per base sequence quality check. As its name suggested, this analysis summarizes over all sequence quality of each sample for each read base. In my case, the read length is 151 base pairs represented in the x-axis. The y-axis on the graph showed the quality scores. The sequence quality is calculated as $Q = -10 \times \log_{10}(e)$ where ‘e’ is the estimated probability of the base call being wrong. Thus, higher score indicates higher quality, and it ranges from 0 to 40. A quality score of 20 represents an error rate of 1 in 100, and a quality score of 40 represents an error rate of 1 in 10,000 and a call accuracy of 99.99%. Green region represents good quality calls from 28 to 40; orange region represents calls of reasonable quality from 20 to 28; red represents poor quality calls with quality less than 20, with a call accuracy of 99%. The quality of calls on most platforms will degrade as the run progresses, so it is common to see base calls falling into the orange area towards the end of a read.

Each column of box and whisker plot is the summarized quality score of all the reads for that particular base position. “The central red line is the median value; the yellow box represents the inter-quartile range (25-75%); the upper and lower whiskers represent the 10% and 90% points. The blue solid line represents the mean quality”[11]. **Figure 2.1C** is the overall per base sequence quality for all sample, which showed 161 passed and 9 samples with warning.

The 9 warning samples are:

- H_VY-61609_S1512483_VI.N.bam
Figure 2.1 Sequence quality check. A) Passed sample from H_VY-1DKYRE_D1202618_I.B.11 and B) Warning sample from H_VY-83775_S1511871_II.G.39i and C) Summary sequence quality score showed 161 passed (green lines) and 9 warning (yellow lines) for DIAN and Knight ADRC dataset.
Per tile sequence quality

Because we used an Illumina library that retains its original sequence identifiers, the sequencing output also documents each read’s flowcell tile information. Thus, for Illumina sequencing data FastQC reports quality scores from each tile across all the base positions to see if there was a loss in quality associated with any particular part of the flowcell. The plot shows the deviation from the average quality for each tile. Cold color indicates good quality and warm color indicates bad quality that a tile had worse qualities than other tiles for that base. A passed sample plot should be blue all over. FastQC user manual explains that “reasons for seeing warnings or errors on this plot could be transient problems such as bubbles going through the flowcell, or they could be more permanent problems such as smudges on the flowcell or debris inside the flowcell lane”[11]. My samples all passed the test except one showed warning (Figure 2.2B). I observed that there was a quality drop on the end of the reads in some tiles of the flowcells.

Figure 2.2 Per tile sequence quality. A) passed sample from H_VY-1DKYRE_D1202618_I.B.11 B) Warning sample from H_VY-62240_S1511620_IV
Per sequence quality scores

The per sequence quality score report allows me to see the overall quality distribution of my reads and to detect if I have reads with universally low-quality scores. If some reads are poorly imaged, for example, when they are on the edge of the field of view, they will have universally poor quality[11]. My samples showed that the majority of reads are good quality ranging from 28 to 40 (Figure 2.3).

Figure 2.3 Per sequence quality scores. A) passed example from H_VY-1DKYRE_D1202618_I.B.11 B) summary results for all samples showed they all passed for this test.
Per Base Sequence Content

Per Base Sequence Content plots out the proportion of DNA bases for each base position, which are the percentage of A, T, C, G. In a random library, I would expect that there would be little to no difference between the different bases of a sequence run, so the lines in this plot should run parallel with each other. The relative amount of each base may not be equally 25% for each type of nucleic acid, but should reflect the overall amount of these bases in the sequenced genome and should not be hugely imbalanced from each other in any case[11].

It is worth noting that some types of library will always produce biased sequence composition, normally at the start of the read, which is what is happening in my samples with Illumina. “Libraries produced by Illumina priming use random hexamers (including nearly all RNA-Seq libraries). Those hexamers were fragmented using transposases, which inherit an intrinsic bias in the positions at which reads start. This bias does not concern an absolute sequence, but instead provides enrichment of a number of different K-mers at the 5' end of the reads. Therefore, it is common for Illumina sequencers to have the first 7-9 bases with unbalanced base sequence contents. While this is a true technical bias, it can be corrected by trimming, but in most cases, it does not seem to adversely affect the downstream analysis”[11]. It will however produce a warning or error in this module when the difference between A and T, or G and C is greater than 10% (warning) and 20% (failed) in any position. After the first nine positions, the lines run flat and in parallel with each other indicating balanced and unbiased nucleic acid contents (Figure 2.4). Due to the technical bias of Illumina library, this module produced 38 warning and 132 failed test results for the dataset.
**Figure 2.4 Per base sequence content.** The percentage of each DNA nucleic acid type for each base position was labeled and color coded respectively.

Per base N content

“If a sequencer is unable to make a base call with sufficient confidence then it will normally substitute an N rather than a conventional base call. This module plots out the percentage of base calls at each position for which an N was called”[11]. My samples only showed a very low N content at the beginning of the reads due to unavoidable library technical bias, and at 100bp for some samples (Figure 2.5). All samples passed this test.
Some high throughput sequencers generate sequence fragments of uniform length, but others can contain reads of wildly varying lengths. Even within uniform length libraries some pipelines will trim sequences to remove poor quality base calls from the end. This module generates a graph showing the distribution of fragment sizes in the

**Figure 2.5 Per base N content.** A) Passed sample from H_VY-1DKYRE_D1202618_I.B.11 and B) summary for all samples.

**Sequence length distribution**

“Some high throughput sequencers generate sequence fragments of uniform length, but others can contain reads of wildly varying lengths. Even within uniform length libraries some pipelines will trim sequences to remove poor quality base calls from the end. This module generates a graph showing the distribution of fragment sizes in the
file which was analyzed"[11]. In many cases this will produce a simple graph showing a peak only at one size, which is the case for my sample shown here (Figure 2.6), but for variable length FastQ files this will show the relative amounts of each different size of sequence fragment[11].

![Sequence Length Distribution]

**Figure 2.6 Sequence length distribution.** Passed sample from H_VY-1DKYRE_D1202618_I.B.11

**Adapter content**

I have adapter contamination due to adapter read-through problem associated with fragmented short reads. Adapter source was predicted and my sample showed potential adapter contamination from Illumina universal adapter (Table 2.4; Figure 2.7A). All samples failed this test (Figure 2.7B).
Expected observations with adapter dimer contamination[11]:

- Drop in per base sequence quality after base 60
- Possible bi-modal distribution of per sequence quality scores
- Distinct pattern observed in per bases sequence content up to base 60
- Spike in per sequence GC content
- Overrepresented sequence matching adapter
- Adapter content > 0% starting at base 1

<table>
<thead>
<tr>
<th>Adapter</th>
<th>i5 index name</th>
<th>i5 index bases</th>
<th>Index</th>
</tr>
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<tbody>
<tr>
<td>D501-D508 adapter</td>
<td>D501</td>
<td>TATAGCCT</td>
<td>D503</td>
</tr>
<tr>
<td></td>
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<td>ATAGAGGC</td>
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<td>D503</td>
<td>CCTATCCT</td>
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<td>GGCTCTGA</td>
<td>D506</td>
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<td></td>
<td>D505</td>
<td>AGGCGAAG</td>
<td>D507</td>
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<td>D508</td>
<td>GTACTGAC</td>
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<th>i7 index bases</th>
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<td></td>
<td>D706</td>
<td>GAATTCTG</td>
</tr>
<tr>
<td></td>
<td>D707</td>
<td>CTGAAAGCT</td>
</tr>
</tbody>
</table>
Sequence duplication level

“The plot shows the proportion of the library which is made up of the duplicated sequences in each of the different duplication level bins. There are two lines on the plot. The blue line takes the full sequence set and shows how its duplication levels are distributed. In the red plot the sequences are de-duplicated and the proportions shown are the proportions of the remained different duplication levels in the original data after removing the duplicated sequences. In a properly diverse library most sequences should fall into the far left of the plot in both the red and blue lines. A general level of
enrichment, indicating broad over sequencing in the library will tend to flatten the lines”[11], lowering the low end and generally raising other categories (Figure 2.8B blue line). More specific enrichments of subsets, or the presence of low complexity contaminants will tend to produce spikes towards the right of the plot (Figure 2.8A blue line). “These high duplication peaks will most often appear in the blue trace as they make up a high proportion of the original library, but usually disappear in the red trace as they make up an insignificant proportion of the deduplicated set. If peaks persist in the red trace then this suggests that there are a large number of different highly duplicated sequences which might indicate either a contaminant set or a very severe technical duplication. The module also calculates an expected overall loss of sequence were the library to be deduplicated shown in the figure headline at the top of the plot, which gives a reasonable impression of the potential overall level of loss”[11].

Note that both biological duplication and technical duplication were not differentiated in this analysis and the way to differentiate these two categories is to examine if the duplicated reads are mostly from physically connected genome regions after alignment[11]. High coverage data has more reads so it is not surprised to see higher duplication level. Notice that among the three passed samples, two are low yield samples. I would also expect the majority of duplications are from rRNA so the samples with high rRNA will have high duplication levels, which is proved by comparing the two figures showing one normal rRNA sample (Figure 2.8A) and one high rRNA sample (Figure 2.8B). In STAR alignment, the default setting for reads aligned to multiple location is 10, and when it’s above 20 it will not map to the reference and those will go to unmapped category, so from the duplication level plot >10 bin percentage I would be able to have a
rough estimation of the percentage of reads that would map to multiple location and unmapped percentage of STAR alignment results. For project that look at unmapped section of the samples this duplication level information might be extremely important, for example circular RNA or microbial RNA focused projects.

Notice that among the three passed samples, two are low yield samples:

- H_VY-12152_S1512495.bam
- H_VY-60410_S1511525_I.D.19.bam (low yield)
- H_VY-62240_S1511620_IV.bam (low yield)

Figure 2.8 Sequence duplication levels. A) Warning sample from H_VY-1DKYRE_D1202618_I.B.11 and B) failed sample from H_VY-82018_S1512310_Il.H.40 and C) summary sample for all samples
Per sequence GC content

“This module measures the GC content across the whole sequence length and compares it to a modelled normal distribution of GC content. In a normal random library, I would expect to see a roughly normal distribution of GC content where the central peak corresponds to the overall GC content of the sequenced genome. An unusually shaped distribution could indicate a contaminated library or some other kinds of biased subset. A normal distribution which is shifted indicates some systematic bias which is independent of base position. If there is a systematic bias which creates a shifted normal distribution then this won't be flagged as an error by the module since software doesn't know what the sequenced genome's GC content should be”[11].

This module will indicate a failure if the sum of the deviations from the normal distribution represents more than 30% of the reads. Warnings in this module usually indicate a problem with the library. Sharp peaks on an otherwise smooth distribution are normally the result of a specific contaminant (adapter dimers for example, Figure 2.9A), which may well be picked up by the overrepresented sequences module. Broader peaks may represent contamination with a different species. In my samples, the distribution showed sharp peaks on an otherwise smooth distribution or severely deviated from normal distribution may indicate adapter dimers contamination, which might be picked by other failed modules, such as overrepresented sequences, adapter content, and kmer content. Overall, I have 42 samples with warning and 128 failed samples (Figure 2.9B).
Overrepresented sequences

“A normal high-throughput library will contain a diverse set of sequences, with no individual sequence making up a tiny fraction of the whole. Finding that a single sequence is very overrepresented in the set either means that it is highly biologically significant, or indicates that the library is contaminated, or not as diverse as expected”[11]. This module lists all of the sequence which make up more than 0.1% of the total (Figure 2.10). “To conserve memory only sequences which appear in the first
100,000 sequences are tracked to the end of the file. For each overrepresented sequence the program will look for matches in a database of common contaminants and will report the best hit it finds. Hits must be at least 20bp in length and have no more than 1 mismatch. Finding a hit does not necessarily mean that this is the source of the contamination, but may point me in the right direction. Because the duplication detection requires an exact sequence match over the whole length of the sequence any reads over 75bp in length are truncated to 50bp for the purposes of this analysis. Even so, longer reads are more likely to contain sequencing errors which will artificially increase the observed diversity and will tend to underrepresent highly duplicated sequences”[11].

### Overrepresented sequences

<table>
<thead>
<tr>
<th>Sequence</th>
<th>Count</th>
<th>Percentage</th>
<th>Possible Source</th>
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<tr>
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<td>CTCAGGCTTGGATGGTGTTCTAGTACCAAGGGATGCTTACTGATTC</td>
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<td>CTCGTTACGGCAGGACTGGTGTTCTCCTGGAGGAGTCACACTAT</td>
<td>194634</td>
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</tr>
</tbody>
</table>

### Figure 2.10 Overrepresented sequences. Warning sample from H_VY-1DKYRE_D1202618_1.B.11.

#### Kmer Content

The Kmer Content module will do a generic analysis of all of the Kmers in my sample library to find those which do not have even coverage through the length of my reads (Figure 2.11). The top six kmers are plotted in the graph. This can find a number of different sources of bias in the library which can include the presence of read-through adapter sequences building up on the end of the sequences. The presence of any
overrepresented sequences in my library (such as adapter dimers) will cause the Kmer plot to be dominated by the Kmers these sequences contain. What I have in my samples are two folds: 1. Unbalanced sequence content for the first 9 bases which is intrinsic technical bias associated with Illumina library. 2. Adapter read through problem at the end of the reads which can be corrected with STAR alignment using adapter soft clipping option.

**Figure 2.11 Kmer Content.** Failed sample from H_VY-1DKYRE_D1202618_I.B.11.
In summary, my DIAN and Knight ADRC samples have good sequence quality in general, reflected in categories such as per base, per tile, per sequence quality scores, and per base N content (Table 2.5). Because we used Illumina sequencing, the first nine bases of each read have technical bias, therefore, none of the samples passed per base sequence content test. However, it does not impact downstream analysis. Due to potential rRNA and adapter contamination, several test metrics captured these observations, for example, per sequence GC content, sequence duplication levels, overrepresented sequences, adapter content, and kmer content. Samples with high rRNA contamination were not different from the other samples in terms of deconvolution results, but will be excluded from other downstream analysis. Adapter sequences could be soft clipped during alignment or trimmed ahead of alignment.

<table>
<thead>
<tr>
<th>Table 2.5 DIAN and Knight ADRC FastQC summary</th>
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<tr>
<td>Pass</td>
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<tr>
<td>-----------------------------------------------</td>
</tr>
<tr>
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<td>Per base sequence quality</td>
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<td>Overrepresented sequences</td>
</tr>
<tr>
<td>Adapter Content</td>
</tr>
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<td>Kmer Content</td>
</tr>
</tbody>
</table>

IGV visualization and IBD to verify sample identity

Aligned and sorted bam files were loaded into IGV[224] to perform visual inspection of target variants. For example, visualization of a PSEN1_S290C delE9 carrier in the top red track is compared to a non-carrier in the bottom blue track shown in Figure.
2.12 using IGV sashimi plot. DelE9 is a mutation (Chr14: 73673093 G>A) in \textit{PSEN1} gene that result in exon 9 exclusion. In this carrier subject, in whom harbors heterozygous exon 9 deletion, the peak in the middle represents gene expression level of \textit{PSEN1} exon 9 flanked by exon 8 and exon 10. Because this carrier is heterozygous for delE9, the exon 9 peak is half in height compared to its neighbors on either side, and there are more reads (79 reads) that skipped exon 9 and linked exon 8 and exon 10 compared to the non-carrier in the bottom track (0 read). Samples carrying unexpected variants or missing expected variants were labeled as potential swapped samples. In addition, variants were called from RNA-Seq following BWA/GATK pipeline\cite{171, 181}. The identity of the samples was later verified by performing IBD analysis against genomic typing from GWAS chipsets.

\textbf{Figure 2.12} \textit{PSEN1} delE9 Sashimi Plot using IGV. Top red track is from sample H\textunderscore VY-2968OM\textunderscore S1512443\textunderscore I.B.13, a PSEN1\textunderscore S290C delE9 carrier, in whom harbors heterozygous exon 9 deletion. Bottom blue track is from a non-carrier subject.
GWAS PCA and Ethnicity Check

GWAS principal component analysis (PCA) components were extracted for matched RNA-Seq subjects to check ethnic identity. Among RNA-Seq subjects there are 6 subjects are African Americans, while the rest are European Americans (Figure 2.13).

![GWAS PCA of ethnicity check](Image)

**Figure 2.13 GWAS PCA of ethnicity check.** HapMap Europeans are color coded as yellow; HapMap Africans are color coded as blue; HapMap Asians are color coded as red. My samples are color coded as black, which mostly fall into the European ethnic group except six subjects labeled with their IDs.
2.3.3 Expression quantification

I applied Salmon transcript expression quantification (ver 0.7.2)[208] to infer the gene expression for all samples included in the reference panel and participants in the Mayo, MSBB, DIAN and Knight-ADRC. I quantified the coding transcripts of *Homo Sapiens* included in the GENCODE reference genome (GRCh37.75). Similarly, I quantified the expression of the mice samples included in the reference panel using the *Mus Musculus* reference genome (mm10).

2.3.4 Reference panel

Reference Samples

I assembled a cell-type specific reference panel from publicly available RNA-Seq datasets comprised of both immunopanning collected or iPSC derived neurons, astrocytes, oligodendrocytes, and microglial cells from human and murine samples. For immunopanning collected cells, antibodies for cell-type specific antigens were utilized to bind and immobilize their targeted cell types in order to immunoprecipitate and purify each cell type from the suspensions[290]. cDNA synthesis was accomplished using Ovation RNA-Seq system V2 (Nugen 7102) and library prepared with Next Ultra RNA-Seq library prep kit from Illumina (NEB E7530) and NEBNext® multiplex oligos from Illumina (NEB E7335 E7500). TruSeq RNA Sample Prep Kit (Illumina) was used to prepare library for paired-end sequence on 100ng of total RNA extracted from each sample. Illumina HiSeq 2000 Sequencer was used to sequence all libraries[290].
Both human adult temporal cortex tissue, collected from patients receiving neurological surgeries, and mice cells were disassociated, sorted and sequenced as described elsewhere[291], and deposited in the Gene Expression Omnibus GSE73721 and GSE52564. I also accessed neural progenitor cells (day 17) and mature human neurons (day 57 and 100) from Broad iPSC deposited in the AMP-AD portal[7] and neural progenitor cells and iPSC-derived neurons from[37]. Broad iPSC derived neurons accessed from AMP-AD portal were generated using an embryoid body-based protocol to differentiate into forebrain neurons[1]. Wild-type cells used in the protocol were obtained from UConn StemCell Core. RNA was purified using PureLink RNA mini-kit (Life Technologies) and libraries were prepared by Broad Institute's Genomics Platform using TruSeq protocol. Please refer to Table 2.6 for additional information.

<table>
<thead>
<tr>
<th>Type</th>
<th>Human</th>
<th>Mouse</th>
<th>Human iPSC(^a)</th>
</tr>
</thead>
<tbody>
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<td>GSM1269905</td>
<td>YZ2-100day</td>
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<tr>
<td></td>
<td></td>
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<tr>
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<td></td>
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<td>GSM1901341</td>
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</tbody>
</table>

\(^a\) Samples accessed from the Broad iPSC cell-lines deposited in the AMP-AD.
Marker Genes

The reference panel was assembled with samples from four distinct cell types. A redundant set of well-known cell-type markers was selected from the literature [41, 131, 291] (Table 2.7). Principal component analysis was performed on the reference panel using R function `prcomp` (version 3.3.3) to verify that the expressions of these gene were clustering samples by their cell types (Figure 2.14b; Figure 2.15a).

Table 2.7 Gene markers for principal brain cell types.

<table>
<thead>
<tr>
<th>Type</th>
<th>Cell Marker</th>
<th>Human</th>
<th>Mouse&lt;sup&gt;a&lt;/sup&gt;</th>
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<td>Syn1</td>
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<td>Astrocyte</td>
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<td></td>
<td>IL1A</td>
<td>Il1a</td>
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</tbody>
</table>

<sup>a</sup> Mouse homologous genes were identified from Mouse Genome Database.
Figure 2.14 PCA of samples included in the reference panel. a) Transcriptome-wide. Genes included in the reference panel b) PC1 vs PC2 and c) PC3 vs PC4.
2.3.5 Inference of the cellular population structure

I ascertained alternative computation deconvolution algorithms implemented in the CellMix package (ver 1.6). Based on accuracy and robustness evaluation results I compared and reported the following three algorithms that outperformed the others: Digital Sorting Algorithm (named “DSA”)[293], which employs linear modeling to infer cell distributions; the method population-specific expression analysis (PSEA, also named meanProfile in CellMix implementation)[157] that calculates estimated expression profiles relative to the average of the marker gene list for each cell type[157]; and a semi-supervised learning method that employs non-negative matrix factorization (ssNMF in CellMix implementation)[103]. I employed a leave-one-out cross-validation procedure to evaluate the accuracy provided by each method. The best performing algorithm ssNMF integrates cell-type marker genes to resolve the drawbacks of completely unsupervised standard non-negative matrix factorization. I followed the standard procedure described in the CellMix package, that included the extraction of marker genes from the reference samples (function extractMarkers from the CellMix package), and the posterior invocation of the function ged to infer cellular population from the gene expression of bulk RNA-Seq data. Besides, I tested additional methods which provided considerably lower accuracy (least-squares fit[8], quadratic programing[108]) or no significant difference (support vector regression[199] or latent variable analysis[51]) to the methods presented.

I selected the reference samples that provide the most faithful transcriptomic profile for their respective cell types by following a leave-one-out cross validation approach. I trained iteratively deconvolution models using all but one of the samples that
was tested. Only samples predicted with a composition higher than 80% were kept for the reference panel (Table 2.6; Figure 2.15b).
Figure 2.15 Leave-one-out evaluation of reference panel. a) Gene expression levels (log-transformed) for reference panel. The cell types of the isolated/iPSC-derived samples are color-coded and labeled on the y-axis. b) A leave-one-out procedure to obtain the cellular proportion for each of the samples of the reference panel was used. Cell-type proportions are shown as stacked percentage (red: astrocytes; green: microglia; blue: neuron; purple: oligodendrocyte).
2.3.6 Accuracy and Robustness Evaluation

Chimeric validation

To emulate heterogeneous tissue with known and controlled cellular composition, I generated chimeric libraries pooling reads (to a total of 400,000) contributed from the human reference samples (See Table 2.6). This process was repeated 720 times, using alternative reference samples to model each cell type. The proportion of reads that the libraries of neurons, astrocytes, oligodendrocytes and microglia provided to the chimeric libraries varied in predefined ranges (Figure 2.16). As a result, each of the chimeric libraries contained reads that followed 32 different distributions (neuronal reads contributed between 2 to 36% of reads, astrocytes between 22 to 76%, oligodendrocytes between 6 to 62% and microglia between 1 to 5%). Refer to Table 2.8 for detailed description of the 32 different distributions. I quantified the chimeric reads using Salmon (v0.7.2)[208], and employed the reference samples that did not contribute reads to the chimeric library as reference panel for the deconvolution methods.

Overall, I applied my digital deconvolution analyses to 23,040 (720 × 32) chimeric libraries. I evaluated the accuracy using the root-mean-square error (RMSE, Equation 2.1) to compare the digital deconvolution cellular proportion estimates (method ssNMF) versus the defined proportion of reads specific to each of the chimeric libraries:

\[
RMSE = \sqrt{\frac{\sum_{i=1}^{n} (\hat{y}_i - y_i)^2}{n}} \quad \text{(Equation 2.1)}
\]

\(\hat{y}_i\) – estimated value, \(y_i\) – observed value
Figure 2.16 Chimeric library deconvolution simulation. Human cell-type specific reference samples contributed 720 different combinations to generate chimeric libraries. Reads were randomly sampled following 32 pre-specified distributions (Neuronal reads contributed between 2 to 36% of reads, astrocytes between 22 to 76%, oligodendrocytes between 6 to 62% and microglia between 1 to 5%). Each chimeric library was quantified and the cellular distribution estimated using digital deconvolution. These estimates were compare to prior distribution.
Table 2.8 Simulated chimeric tissue cell-type composition. Percentages of reads contributed to the synthetic chimeric libraries.

<table>
<thead>
<tr>
<th>Configurations</th>
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<th>Astrocyte</th>
<th>Oligodendrocyte</th>
<th>Microglia</th>
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<td>21.3</td>
<td>60.6</td>
<td>13.5</td>
<td>4.57</td>
</tr>
</tbody>
</table>

For each target cell type the distribution was pre-defined to cover a broad range of biological viable proportions (highlighted in bold).
I also tested whether the deconvolution results were dominated by the expression of any specific marker gene and ascertained the robustness of the inferred cellular population structure to any possibly altered expression of marker genes. To do so, I performed the deconvolution analysis discarding each of the marker genes one at a time and evaluated how these distributions differed in comparison to the full gene reference panel.

### 2.3.7 Statistical Analysis

**Transcriptome PCA and covariate analysis**

Number of reads gene quantification results from Salmon were normalized with DeSeq2’s VarianceStabilizingTransformation function after removing genes with total reads count less than 10,000. First 10 principle components were extracted from PCA, and single and stepwise covariate analysis using linear model were performed to investigate what covariates would affect data quality and downstream analysis. RNA quality (RIN and DV200), post-mortem index (PMI), sequencing pooling, sex, age at death, and brain tissue origin were included in either single covariate and stepwise covariate analysis. The results showed RNA quality measurements RIN and DV200, and age at death are the most important confounding factors that impact the analysis. Covariate correlation was performed to examine any correlation among the factors to avoid including highly correlated covariates that lead to over correction of the model, for example, RIN and DV200 are associated with quality but they are highly correlated as well. Since RIN and DV200 are highly correlated, using RIN only is recommended because other publicly ascertainment datasets only have RIN measurements. (Figure 2.17). Later in Chapter 3, I will replace RIN for transcript integrity number (TIN), which will be inferred directly from RNA-Seq data. Notice that ribosomal RNA contents are
negatively correlated with uniquely mapped reads, because those rRNA will be mapped to multiple locations due to its highly conserved and similar sequences. Besides, uniquely mapped reads also anti-correlated with incorrect strand reads, percentage of reads that mapped to multiple loci, and median 5’ to 3’ bias. The neuronal and astrocyte proportion I inferred from the RNA-Seq data are also highly negatively correlated.

**Figure 2.17 Covariate correlation.** Major covariates of RNA-Seq and inferred cell type compositions from deconvolution were correlated to avoid including highly correlated covariates in downstream analysis.
Transcriptome-wise PCA showed seven subjects are very different from the rest and they appeared as outlier on PC1 vs PC2 plot (Figure 2.18). iPSC samples were excluded from this analysis and analyzed separately in another study[143]. Among the seven outliers, 60007 and 83774 also have high median 5’ to 3’ bias. The top 30 genes that contributed the most to PC1 and top 30 to PC2 were extracted and plotted as a heatmap to show the dramatic difference between these outlier subjects and the rest of the samples (Figure 2.19). Notice that the left most 5 subjects are the outliers on transcriptome-wise PCA, but the other two outliers 83774 and 60007 do not have an obvious clustering pattern in the heatmap, suggesting that high median 5’ to 3’ bias also contribute variances observed in PC1 and PC2.

Figure 2.18 Transcriptome-wise principle component analysis. Subject transcriptome-wise PCA were plotted using PC1 and PC2 with outliers labeled with subject ID.
Figure 2.19 Top 60 Genes for PC1 and PC2 heatmap. The left most 5 subjects are the outliers on transcriptome-wise PCA, and they are clustered as a separate group in the top 30 genes contributing the most variance to PC1 and PC2. The other two outliers 83774 and 60007 also have high median 5’ to 3’ bias that do not have an obvious clustering pattern in the graph.
Cell type proportions and disease status association analysis

I employed linear regression models to test the association between cell-type proportions and disease status (R Foundation for Statistical Computing, ver. 3.3.3). Stepwise discriminant analysis (stepAIC function of R package MASS, version 7.3-45) was used to determine significant covariates, and correct for confounding effects. I included RNA integrity number (RIN), batch, age at death and post-mortem interval (PMI) as covariates for the Mayo Clinic analyses. For Mount Sinai Brain Bank analyses, I corrected for RIN, PMI, race, batch and age at death. I also used linear-mixed models to perform multiple-region association analysis, employing random slopes and random intercepts grouping by observations and by donors[253], and correcting for the same covariates previously described.

To analyze the DIAN and Knight-ADRC studies I applied linear-mixed models (function lmer and Anova, R packages lme4 ver. 1.1 and car ver. 2.1, respectively), clustering at family level to ascertain the effect of the neuropathological status in the cell proportion, and corrected for RIN and PMI. For late-onset specific analyses I also corrected for age at death. Cellular composition shown as proportions were plotted using R package ggplot2 (ver 2.2.1).

2.4 Results

2.4.1 Study design

To infer cellular composition from RNA-Seq, I firstly assembled a reference panel to model the transcriptomic signature of neurons, astrocytes, oligodendrocytes and microglia. The panel was created by analyzing expression data from purified cell lines. I
evaluated alternative digital deconvolution methods and selected the best performing for my primary analyses. I tested the digital deconvolution accuracy on induced pluripotent stem cell (iPSC) derived neurons/microglia cells and neuronal Translating Ribosome Affinity Purification followed by RNA-Seq (TRAP-seq; Figure 2.20). Finally, I verified its accuracy by creating artificial admixture with pre-defined cellular proportions.

**Figure 2.20 Study design.** Development of the brain cell-type transcriptomic reference panel (left column): the expression signatures of key cell types of the brain were curated by compiling publicly available RNA-Seq data from neurons, astrocytes, oligodendrocytes and microglia. The panel was curated iteratively to retain only those samples that showed the most faithful expression signature, while evaluating alternative digital deconvolution methods. The accuracy of digital deconvolution to estimate brain cellular proportion was validated using additional cell-type specific samples, and also by generating chimeric libraries. To study cellular population structure in AD (right column), I accessed publicly available datasets from the Advanced Medicines Partnership-AD knowledge portal (AMP-AD), including Mayo Clinic and Mount Sinai Brain Bank datasets. In addition, we generated RNA-Seq from participants of the Knight-ADRC and The Dominantly Inherited Alzheimer (DIAN) studies. These three studies generated RNA-Seq data from pathological aging brains, Alzheimer’s disease cases, and neuropath-free controls for a total of six cerebral cortex regions and cerebellum. I quantified the gene expression for all of the samples included in these studies using the same RNA-Seq processing pipeline. Using digital deconvolution methods, I estimated the brain cellular proportions of the samples and compared the proportion between AD cases and controls. I study the cell structure of brains carriers of Mendelian pathological mutations and variants that confer high-risk to AD. Anterior prefrontal cortex – APC; superior temporal gyrus – STG; parahippocampal gyrus – PHG; inferior frontal gyrus – IFG; Mount Sinai Brain Bank – MSBB; Alzheimer’s disease – AD; pathological aging – PA.
Once the deconvolution approach was optimized, I calculated the cell proportion in AD cases and controls from the different brain regions of Mayo and MSBB datasets. The RNA-Seq data for the Mayo Clinic study (N = 191)[9] and Mount Sinai (MSSM) Brain Bank (MSBB; N = 300)[3] are deposited in the Advanced Medicines Partnership-AD (AMP-AD) knowledge portal (Synapse ID: syn5550404 and syn3157743; Table 2.1). The Mayo study includes RNA-Seq from the temporal cortex and cerebellum for AD affected and non-demented controls, in addition to pathological aging participants (Figure 2.20). The MSBB also profiled four additional cerebral cortex areas: anterior prefrontal cortex - APC, superior temporal gyrus - STG, parahippocampal gyrus – PHG, and inferior frontal gyrus – IFG; Table 2.1; Figure 2.20). I restricted the case-control analysis to subjects with definite AD and autopsy confirmed controls. In addition, we generated RNA-Seq from parietal lobe for participants of the Knight-ADRC (84 late-onset cases, carriers of genetic risk factors and 16 controls; Table 2.2) and The Dominantly Inherited Alzheimer Network (DIAN; 19 carriers of mutations in APP, PSEN1, PSEN2) (Table 2.1; Figure 2.20). I employed the same pipeline to process all of the samples in order to avoid any bias. Furthermore, RNA-Seq from the Knight-ADRC and DIAN studies allowed us to compare the cell composition from ADAD vs LOAD brains, and similarly to test for differences in brain of controls, sporadic AD who do not carry any known high-risk variant, carriers of high-risk variants in TREM2 (N = 20), PLD3 (N = 33), and APOE ε4 allele.
2.4.2 Development of a reference panel to estimate brain cellular population structure

Due to limited availability of brain cell-type specific transcriptomic data, I compiled reference samples from different sources, including single-population RNA-Seq from mice and human (immunopan-purified oligodendrocytes, neurons, astrocytes and microglia and iPSC-derived neurons and astrocytes).

I selected 17 well accepted genes that tag brain cell types based on literature reviews[41, 131, 291]. A visual inspection of the expression of these marker genes in the samples I compiled suggested a divergent transcriptomic profile among the cell types (Figure 2.15a). The PCA showed that their expression was sufficient to cluster samples of neurons, astrocytes, oligodendrocytes and microglia with their respective cell types, regardless of the species of the reference samples (Figure 2.14b; Table 2.6). I observed that first principal component (PC) captured the expression profile of astrocytes; as shown by the significant association of the expression of astrocyte marker genes with (p < 8.05×10^{-03}). The second PC captured the expression of genes whose expression is characteristic to oligodendrocytes (p < 2.52×10^{-02}). The third PC was negatively associated with neuronal genes (p < 1.11×10^{-05}) and positively with microglia (p < 1.42×10^{-02}). Overall, the principal component analysis (Figure 2.14b) indicated that these genes can effectively cluster samples by their cell-type.

Given the technical and biological heterogeneity of the samples I compiled for reference panel, I carried out an optimization phase to identify those samples that showed the most faithful expression profile to represent their respective cell types (See Methods). From the leave-one-out cross-validation results, I noticed that not all of the
cell-type specific samples were predicted as expected (defined with a correct prediction proportion higher than 80%). Samples failed this criteria were due to various reasons, for example, the expression profile of immunopan-purified astrocytes collected from mice[291], human fetal[291] or human sclerotic hippocampal[291] brains were reported with altered expression[38, 291] that differed to an extent that could not be accurately ascertained by deconvolution methods. Similarly, neuronal proportion inferred from iPSC-derived neurons from schizophrenic donors[38] and iPSC-derived neurons collected at early stages of differentiation (< 100 days; Synapse ID: syn3607401) were also lower than 80%. These samples did not cluster with their expected cell types in the marker gene PCA either, and coincidently the leave-one-out cross-validation indicated that these samples had an expression signature that differed from the other samples of the same cell type.

To evaluate the performance of reference panel performance and test out different deconvolution algorithms, I employed and compared six digital deconvolution methods implemented in the CellMix package (ver 1.6) to infer cellular composition from reference samples RNA-Seq data, including qprog[108], cs-qprog, DSA[293], ssFrobenius[103], meanProfile[157], deconf[221]. The deconvolution performance of reference panel was evaluated by following a leave-one-out cross-validation procedure to compare the predicted cellular composition with its expected cellular identity of each cell-type specific sample. The accuracy of this comparison was quantified using the root-mean-squared error (RMSE) calculation. A semi-supervised method adapted from non-negative matrix factorization[103] (ssNMF – named ssFrobenius in CellMix) generated the most accurate predictions; and I verified that similar results were obtained by the

I ascertained the effect that sequencing depth has in the accuracy of deconvolution. I generated low-coverage versions (Picard DownsampleSam ver 2.8.2) of the samples that included a reduced number of randomly sampled reads (400,000 reads per sample), quantified the gene expression, inferred their cellular population proportions, and compared the distribution estimates with their full-depth libraries sequencing (more than 30 million reads per sample). I observed that the deconvolution was robust to the sequencing coverage, as shown by a correlation \( r^2 = 0.98 \) (\( p < 2.2 \times 10^{-16} \); Figure 2.21). My final reference panel (Table 2.6; Table 2.7) had a very high confidence to predict cell types with a mean predicted accuracy = 95.2%; s.d. = 4.3, and a root-mean-square error (RMSE) = 0.06 (Table 2.9).
Figure 2.21 Comparison of cell proportions estimated from full-depth and downsampled RNA-Seq data. Each sample of the reference panel sample was down-sampled (400,000 reads) and cellular population structure inferred following leave-one-out procedure. Cell-type proportions of the samples inferred using the full-depth RNA-Seq data are presented along the X-axis, and along the Y-axis the counterparts inferred using shallow RNA-Seq.
Table 2.9 Evaluation of deconvolution accuracy. Overall and cell-type specific root-mean-squared error (RMSE) for reference panel, calculated using the leave-one-out approach for three deconvolution algorithms implemented in CellMix package.

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>Overall</th>
<th>Neuron</th>
<th>Astrocyte</th>
<th>Oligodendrocyte</th>
<th>Microglia</th>
</tr>
</thead>
<tbody>
<tr>
<td>ssNMF&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.064</td>
<td>0.054</td>
<td>0.055</td>
<td>0.028</td>
<td>0.017</td>
</tr>
<tr>
<td>PSEA&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.089</td>
<td>0.08</td>
<td>0.052</td>
<td>0.058</td>
<td>0.025</td>
</tr>
<tr>
<td>DSA&lt;sup&gt;c&lt;/sup&gt;</td>
<td>0.465</td>
<td>0.32</td>
<td>0.328</td>
<td>0.291</td>
<td>0.295</td>
</tr>
</tbody>
</table>

<sup>a</sup> ssNMF: semi-supervised learning non-negative matrix factorization.

<sup>b</sup> PSEA: population-specific expression analysis (also named meanProfile in CellMix implementation).

<sup>c</sup> DSA: Digital sorting algorithm.
2.4.3 Optimization, validation and accuracy estimation of the reference panel and digital deconvolution method

Once I identified the optimal approach to perform digital deconvolution from brain RNA-Seq, I benchmarked it by using three sets of independent pure cell populations and simulated chimeric libraries.

I firstly validated the accuracy to predict neuronal composition by generating RNA-Seq for eight iPSC-derived cortical neurons (see Methods). I observed an accurate prediction in these independent cell lines (mean neuronal proportion = 94.8% and s.d. = 1.1%; Figure 2.22a). I also ascertained the cellular composition of mRNA extracted from the barrel cortex neurons isolated by Translating Ribosome Affinity Purification (TRAP) in 24 mice. TRAP is a method that captures cell-type specific mRNA translation by purifying tagged ribosomal subunit and capturing the mRNA it bound to[122]. I observed an average of neuronal proportion = 96.7% and s.d. = 1.2% (Figure 2.22b).

Similarly, I assessed the RNA-Seq data generated for iPSC-derived microglia (N = 10) deposited in the AMP-AD portal (Synapse ID: syn7203233) and inferred their cellular population structure and observed a mean microglia proportion = 86.6% and s.d. = 7.1% (Figure 2.22c).
Figure 2.22 Cellular population structure of cell-type specific samples. Cell-type proportions shown as stacked percentage (red: astrocytes; green: microglia; blue: neuron; purple: oligodendrocyte). a) iPSC derived cortical neurons (N = 8). b) mouse barrel cortex neurons isolated by Translating Ribosome Affinity Purification (TRAP) procedure (N = 24). c) iPSC derived microglia (N = 10).
To evaluate the accuracy of digital deconvolution for measuring cell-type proportion from cell-type admixtures, I simulated RNA-Seq libraries by pooling reads from individual cell types into well-defined proportions. I combined randomly sampled reads from neurons, astrocytes, oligodendrocytes and microglia to create chimeric libraries that mimic bulk RNA-Seq from brain, but with a range of pre-defined cell-type distributions (Figure 2.16). I then quantified the gene expression for the chimeric libraries and inferred the cell-type distribution (employing for the reference panel samples that did not contribute reads to the chimeric libraries). This process was repeated 23,040 times, choosing distinct human samples to represent each cell type and varying the proportions in 32 alternative distributions (See methods and Table 2.8). The overall error (RMSE) compared to known proportions = 0.08.

Finally, I evaluated whether any gene included in the reference panel was dominating the inference of cell proportions. I re-calculated the cell-type distributions of the chimeric libraries, but dropping each of the genes from the reference panel one at a time. I observed a negligible difference between the cellular population structure inferred using the full reference and the gene-dropped panels (average RMSE = 0.022, s.d. < 0.01). In this way, I verified that the proportions inferred using the reference panel are not driven by the expression of a single gene. This reassured us the inference should be robust to any bias introduced by the potential association of a single gene included in the reference panel with a particular trait.
2.4.4 Deconvolution of bulk RNA-Seq of non-demented and AD brains shows a characteristic signature for neurodegeneration

Pathologically, AD is associated with neuronal death and gliosis specifically in the cerebral cortex. I evaluated whether I could exploit deconvolution methods using my reference panel to detect altered cellular population structure from the bulk RNA-Seq, and whether this corresponded to known pathological alterations.

I initially analyzed the RNA-Seq from the Mayo Clinic Brain Bank that includes bulk RNA-Seq from the temporal cortex (TC) and cerebellum (CB) for 191 participants[9] (Table 2.1). In the TC, I observed a significant higher astrocyte relative proportion ($\beta = 0.23; p = 5.01 \times 10^{-09}$; Table 2.10; Figure 2.25; Table 2.11) in AD brains compared to controls brains. I also found a significant lower relative proportion of neurons ($\beta = -0.17; p = 1.58 \times 10^{-07}$; Table 2.10; Figure 2.25; Table 2.11) and oligodendrocytes ($\beta = -0.07; p = 1.8 \times 10^{-02}$; Table 2.10; Figure 2.23; Table 2.11). As expected, given the absence of pathology, I did not observe a significant difference in the cell-type composition in the CB (Table 2.10).
<table>
<thead>
<tr>
<th>Brain Regions</th>
<th>Sample Size</th>
<th>Neuron</th>
<th>P-value</th>
<th>Astrocyte</th>
<th>P-value</th>
<th>Oligodendrocyte</th>
<th>P-value</th>
<th>Microglia</th>
<th>P-value</th>
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<td></td>
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<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cerebellum</td>
<td>119</td>
<td>-0.03</td>
<td>2.74×10⁻⁰¹</td>
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<td>8.65×10⁻⁰²</td>
<td>-0.02</td>
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<td>-3.19×10⁻⁰⁴</td>
<td>9.19×10⁻⁰⁴</td>
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<tr>
<td>Temporal Cortex</td>
<td>119</td>
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<td>0.23</td>
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<td></td>
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<tr>
<td>Anterior Prefrontal Cortex</td>
<td>184</td>
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<td>8.11×10⁻⁰⁵</td>
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<td>-3.18×10⁻⁰³</td>
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<td>1.35×10⁻⁰¹</td>
</tr>
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<td>-0.04</td>
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<td>Braak Staging</td>
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<tr>
<td>Anterior Prefrontal Cortex</td>
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<tr>
<td>Superior Temporal Gyrus</td>
<td>158</td>
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<td>2.22×10⁻⁰⁷</td>
<td>0.02</td>
<td>2.77×10⁻⁰⁷</td>
<td>-2.91×10⁻⁰³</td>
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<td>-5.47×10⁻⁰⁴</td>
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<td>2.37×10⁻⁰²</td>
<td>-1.01×10⁻⁰³</td>
<td>1.74×10⁻⁰²</td>
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<tr>
<td>Mean Amyloid Plaques</td>
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<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anterior Prefrontal Cortex</td>
<td>184</td>
<td>-1.88×10⁻⁰³</td>
<td>3.6×10⁻⁰³</td>
<td>2.82×10⁻⁰³</td>
<td>1.03×10⁻⁰⁴</td>
<td>-7.99×10⁻⁰⁴</td>
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<td>0.01</td>
<td>4.63×10⁻⁰⁸</td>
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<td>9.01×10⁻⁰²</td>
<td>-2.04×10⁻⁰⁴</td>
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<tr>
<td>Parahippocampal Gyrus</td>
<td>160</td>
<td>-4.96×10⁻⁰³</td>
<td>5.05×10⁻⁰⁹</td>
<td>0.01</td>
<td>1.26×10⁻¹⁰</td>
<td>-9.99×10⁻⁰⁴</td>
<td>1.85×10⁻⁰³</td>
<td>-2.1×10⁻⁰⁴</td>
<td>2.58×10⁻⁰²</td>
</tr>
<tr>
<td>Inferior Frontal Gyrus</td>
<td>159</td>
<td>-2.58×10⁻⁰³</td>
<td>3.82×10⁻⁰⁴</td>
<td>3.53×10⁻⁰³</td>
<td>1.96×10⁻⁰⁵</td>
<td>-7.41×10⁻⁰⁴</td>
<td>1.51×10⁻⁰²</td>
<td>-2.04×10⁻⁰⁴</td>
<td>1.26×10⁻⁰²</td>
</tr>
</tbody>
</table>

The cell-type proportions from AD cases and control were inferred from bulk RNA-seq using the ssNMF method. Effects of AD and associations with additional clinical and pathological phenotypes in cell-type distributions were estimated using linear regression model.
The distribution of microglia was similar in the TC and CB from AD and control brains (Table 2.10; Figure 2.23). The proportion of microglia was lower than any other cell types. The Mayo dataset also includes brains from individuals with pathological aging (PA; Table 2.1); which is neuropathologically defined by amyloid β (Aβ) senile plaque deposits but little or no neurofibrillary tau pathology[9, 192]. I observed a significant lower relative proportion of microglia in PA brains compared to AD in both TC and CB (Table 2.12; Figure 2.24)[169]. Therefore, I speculated that the lack of changes in the AD microglial population was neither due to low statistical power nor the inability of my method to estimate the microglial proportions, but reflected unaltered neuropathological observations in AD brains.

Figure 2.23 Microglia and oligodendrocyte proportions inferred from RNA-Seq of Mayo Clinic and Mount Sinai Brain Bank (MSBB) studies. Mean microglial (green) and oligodendrocyte (purple) proportion for AD cases and neuropath-free controls (bars indicate the standard deviation). The numbers of subjects are indicated below x-axis.
Figure 2.24 Cellular population structure for Alzheimer’s disease (AD) and Pathological Aging (PA) subjects included in the Mayo Clinic study. Columns height represent the mean proportions. The numbers of subjects for each group is reported below x-axis.
I also analyzed data from the MSBB, which contains bulk RNA-Seq for four additional cerebral cortex areas (APC, STG, PHG, IFG). Replicating my findings from the Mayo dataset I observed a significant lower relative proportion in neurons and increase in astrocytes in all four areas (Table 2.10; Figure 2.25; and Table 2.11). The strongest effect size was detected in the parahippocampal gyrus and superior temporal gyrus (p < 3.49×10⁻⁰⁷) (Table 2.10; Table 2.13). Neuropathological studies have described that the parahippocampal gyrus in one of the first brain areas in which AD pathology occurs [33, 78, 267]. I also observed a significant and strong correlation between neuronal and astrocyte relative proportions and last ascertained clinical status (Clinical Dementia Rating - CDR), and number of amyloid plaques and Braak staging (Table 2.10; Figure 2.25; Figure 2.26).
Figure 2.25 Cell-type distributions of the samples included in the Mayo Clinic and Mount Sinai Brain Bank. Mean neuronal (blue) and astrocytic proportion (red) for a) Alzheimer’s disease affected brains (AD) and controls (bars indicate standard deviations). The numbers of subjects for each group are shown below the x-axis. Distribution for additional clinical and pathological phenotypes reported for the Mount Sinai Brain Bank (MSBB): b) clinical dementia rating scores (CDR) and c) Braak and Braak staging. d) Brain cell-type proportions (x-axis) plotted against the mean number of amyloid plaque (values greater than 0; y-axis). Standard errors were depicted in shaded area with LOESS smooth curve fitted to cell-type proportions derived from deconvolution. (** P< 0.01; *** P< 1.0×10^{-3}; and **** P< 1.0×10^{-4}).
Figure 2.26 Neurons and astrocytes distributions for the brains included in the Mount Sinai Brain Bank stratified by CDR and Braak staging. Neuron (blue) and astrocyte (red) proportions for the plotted against a) CDR. b) Braak Staging.
Table 2.11 Comparison of the cellular proportions estimated using the method PSEA in AD and control brains from the Mayo and Mount Sinai Brain Bank.

<table>
<thead>
<tr>
<th>Brain Regions</th>
<th>Sample Size</th>
<th>Neuron</th>
<th>Astrocyte</th>
<th>Oligodendrocyte</th>
<th>Microglia</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mayo</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cerebellum</td>
<td>119</td>
<td>-0.04</td>
<td>0.06</td>
<td>0.02</td>
<td>0.07</td>
</tr>
<tr>
<td>Temporal Cortex</td>
<td>119</td>
<td>-0.17</td>
<td>0.23</td>
<td>3.1×10^{-02}</td>
<td>3.91×10^{-04}</td>
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<td>Mount Sinai Brain Bank</td>
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<tr>
<td>Anterior Prefrontal Cortex</td>
<td>184</td>
<td>-0.04</td>
<td>0.06</td>
<td>0.01</td>
<td>0.01</td>
</tr>
<tr>
<td>Superior Temporal Gyrus</td>
<td>167</td>
<td>-0.07</td>
<td>0.09</td>
<td>8.55×10^{-02}</td>
<td>1.32×10^{-01}</td>
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<td>160</td>
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<td>2.44×10^{-01}</td>
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<tr>
<td>Inferior Frontal Gyrus</td>
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<td>-0.04</td>
<td>0.06</td>
<td>1.15×10^{-01}</td>
<td>1.84×10^{-02}</td>
</tr>
</tbody>
</table>

Table 2.12 Cell-type proportions comparison of subjects diagnosed with Pathological Aging. The cell-type proportions inferred from RNA-seq data using the ssNMF method. Distribution in Pathological Aging (PA) brains, AD cases and neuropath-free controls are compared using linear regression model.

<table>
<thead>
<tr>
<th>Sample Size</th>
<th>Neuron</th>
<th>Astrocyte</th>
<th>Oligodendrocyte</th>
<th>Microglia</th>
</tr>
</thead>
<tbody>
<tr>
<td>PA vs AD</td>
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<tr>
<td>PA vs Control</td>
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<td>Temporal Cor</td>
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<td>-0.05</td>
<td>-0.03</td>
<td>2.75×10^{-02}</td>
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</table>
The cellular population structure differs between ADAD vs LOAD

While the loss of neurons is a common feature of AD, it is not clear whether the mechanism holds true across different forms of AD or AD cases carrying different genetic risk variants. Therefore, I investigated whether AD with distinct etiologies showed different cellular compositions. We generated RNA-Seq data from the parietal lobe of participants enrolled in Knight-ADRC (84 LOAD, 3 ADAD, and 16 neuropath-free controls) and DIAN (19 ADAD) studies (Table 2.1; Table 2.2). I selected the LOAD and ADAD participants to match for CDR at death, brain weight and sex distributions (See Table 2.2).

Using digital deconvolution, I determined the cellular composition for these brains. I observed a significant lower relative proportion of neurons ($\beta = -0.02, p = 2.66\times10^{-02}$) and significant higher relative proportion of astrocyte in AD ($\beta = 0.03, p = $...
5.48×10^{-03}) for the combined LOAD and ADAD brains compared to controls (Table 2.14; Figure 2.27; Table 2.15), consistent with my findings in the Mayo and MSBB datasets. Similarly, the joint analysis of the brains from Knight-ADRC and DIAN showed a significant association between the neuronal and astrocyte relative proportions and neuropathological measures (Braak staging: β = -0.03, p = 8.51×10^{-06} for neurons and β = 0.03, p = 3.83×10^{-06} for astrocytes; Table 2.14; Figure 2.27b) as well as for clinical measures (CDR: β = -0.02, p = 2.66×10^{-02} for neurons and β = 0.03 and p = 5.48×10^{-03} for astrocytes; Table 2.14; Figure 2.27c). I did not observe a significant difference in the compositions of microglia or oligodendrocytes (Table 2.14; Fig S8).
Table 2.14 Cellular population structure altered in the parietal lobe from AD brains in the DIAN study and Knight-ADRC brain bank.

<table>
<thead>
<tr>
<th>Disease Status</th>
<th>Sample Size</th>
<th>Neuron</th>
<th>Effect</th>
<th>P-value</th>
<th>Astrocyte</th>
<th>Effect</th>
<th>P-value</th>
<th>Oligodendrocyte</th>
<th>Effect</th>
<th>P-value</th>
<th>Microglia</th>
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<tr>
<td>AD(^a) vs Control</td>
<td>122</td>
<td>-0.11</td>
<td>5.52×10(^{-4})</td>
<td>0.14</td>
<td>2.48×10(^{-5})</td>
<td>-0.03</td>
<td>6.5×10(^{-2})</td>
<td>-2.64×10(^{-3})</td>
<td>2.49×10(^{-1})</td>
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<tr>
<td>ADAD vs Control</td>
<td>38</td>
<td>-0.19</td>
<td>3.94×10(^{-7})</td>
<td>0.24</td>
<td>1.57×10(^{-10})</td>
<td>-0.04</td>
<td>8.5×10(^{-3})</td>
<td>-0.01</td>
<td>7.77×10(^{-5})</td>
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<tr>
<td>LOAD vs Control</td>
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<td>0.12</td>
<td>3.34×10(^{-4})</td>
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<td>4.57×10(^{-1})</td>
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<td>ADAD vs LOAD</td>
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<tr>
<td>Braak matched</td>
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<td>1.03×10(^{-2})</td>
<td>0.11</td>
<td>9.26×10(^{-4})</td>
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<td>Braak corrected</td>
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<td>4.71×10(^{-3})</td>
<td>0.11</td>
<td>5.24×10(^{-4})</td>
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<td>1.77×10(^{-1})</td>
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<tr>
<td>AD(^a) and Controls</td>
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<td>-0.02</td>
<td>2.66×10(^{-2})</td>
<td>0.03</td>
<td>5.48×10(^{-3})</td>
<td>-0.01</td>
<td>2×10(^{-1})</td>
<td>-4.63×10(^{-4})</td>
<td>4.77×10(^{-1})</td>
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<td><strong>Braak Staging</strong></td>
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<tr>
<td>AD(^a) and Controls</td>
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<td>8.51×10(^{-6})</td>
<td>0.03</td>
<td>3.83×10(^{-6})</td>
<td>-4.24×10(^{-3})</td>
<td>2.04×10(^{-1})</td>
<td>-2.52×10(^{-4})</td>
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<tr>
<td>ADAD and Controls</td>
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<td>2.37×10(^{-5})</td>
<td>0.06</td>
<td>2.45×10(^{-5})</td>
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<td>2.29×10(^{-1})</td>
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<td>4.89×10(^{-1})</td>
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<tr>
<td>LOAD and Controls</td>
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<td>-0.03</td>
<td>7.41×10(^{-4})</td>
<td>0.03</td>
<td>4.63×10(^{-4})</td>
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</tbody>
</table>

\(^a\) AD includes both autosomal dominant AD (ADAD) and late-onset AD (LOAD).

The cellular population structure was inferred using the ssNMF method. Effects and p-values for the association with disease status, clinical dementia rating and Braak staging using generalized mixed models. We identified similar trends with approximately the same significance levels.
Table 2.15 Comparison of the cellular proportions estimated using the method PSEA in AD and control brains from the DIAN and Knight-ADRC

<table>
<thead>
<tr>
<th>Disease Status</th>
<th>Sample Size</th>
<th>Neuron</th>
<th>Effect</th>
<th>P-value</th>
<th>Astrocyte</th>
<th>Effect</th>
<th>P-value</th>
<th>Oligodendrocyte</th>
<th>Effect</th>
<th>P-value</th>
<th>Microglia</th>
</tr>
</thead>
<tbody>
<tr>
<td>AD\textsuperscript{a} vs Control</td>
<td>122</td>
<td>-0.1</td>
<td>1.41×10\textsuperscript{-03}</td>
<td>0.12</td>
<td>7×10\textsuperscript{-05}</td>
<td>-0.02</td>
<td>1.97×10\textsuperscript{-01}</td>
<td>-1.84×10\textsuperscript{-03}</td>
<td>2.4×10\textsuperscript{-01}</td>
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<td>ADAD\textsuperscript{b} vs Control</td>
<td>38</td>
<td>-0.18</td>
<td>2.35×10\textsuperscript{-06}</td>
<td>0.22</td>
<td>3.03×10\textsuperscript{-10}</td>
<td>-0.05</td>
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<td>-3.77×10\textsuperscript{-03}</td>
<td>1.56×10\textsuperscript{-04}</td>
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<tr>
<td>LOAD\textsuperscript{c} vs Control</td>
<td>100</td>
<td>-0.08</td>
<td>1.16×10\textsuperscript{-02}</td>
<td>0.1</td>
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<td>-0.02</td>
<td>2.5×10\textsuperscript{-01}</td>
<td>-1.12×10\textsuperscript{-03}</td>
<td>4.91×10\textsuperscript{-01}</td>
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<td></td>
</tr>
<tr>
<td>ADAD vs LOAD</td>
<td>106</td>
<td>-0.09</td>
<td>9.47×10\textsuperscript{-04}</td>
<td>0.12</td>
<td>1.1×10\textsuperscript{-05}</td>
<td>-0.03</td>
<td>6.23×10\textsuperscript{-02}</td>
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<td>9.38×10\textsuperscript{-01}</td>
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</tbody>
</table>

\textsuperscript{a} AD Includes all of the AD affected subjects from the Knight-ADRC and DIAN studies.

\textsuperscript{b} ADAD: autosomal dominant AD, carriers of pathogenic mutation in \textit{APP}, \textit{PSEN1} or \textit{PSEN2}.

\textsuperscript{c} LOAD: late-onset AD patients not carrying any autosomal dominant mutation.
Figure 2.27 Neuron and astrocyte distributions from the DIAN and Knight-ADRC brains. a) Mean neuronal (blue) and astrocytic (red) proportions for carriers of pathogenic mutations in APP, PSEN1 or PSEN2 (ADAD), late-onset AD (LOAD) and neuropath-free controls (bars indicate standard deviations). Neuronal and astrocytic proportions plotted against b) Braak Staging; c) by Clinical Dementia Rating. d) Cell-type distributions for carriers of AD genetic risk factors. Lines indicate significance levels (*P< 0.05; ** P< 0.01; *** P< 1.0×10⁻³, **** P< 1.0×10⁻⁴).
Next, I compared the cell proportion of LOAD vs ADAD and found that the cell composition differs between them. I firstly selected the LOAD brains (N = 25) to match the Braak staging distribution of ADAD brains (N = 17). The ADAD brains showed a significant lower neuronal proportion compared to LOAD brains (β = -0.08; p = 1.03×10^{-02}; Table 2.14), and increased relative astrocyte proportion (β = 0.11; p = 9.26×10^{-04}; Table 2.14). Then, I analyzed the entire Knight-ADRC LOAD brains, by extending the model to correct for Braak stages. I also observed significant lower relative neuronal proportion (β = -0.09; p = 4.71×10^{-03}; Table 2.14; Figure 2.27a; Table 2.15) and increased relative astrocyte proportion (β = 0.11; p = 5.24×10^{-04}; Table 2.14; Figure 2.27a; Table 2.15) in ADAD brains compared to LOAD. I observed the same cellular differences when I corrected for CDR at death (β = -0.12; p = 2.11×10^{-03} for neurons and β = 0.13; p = 6.29×10^{-04} for astrocytes; Table 2.14; Figure 2.27bc). In summary, my results indicate that ADAD individuals present a higher neuronal death even in the same stage of the disease, suggesting that in ADAD neuronal death play a more important role in pathogenesis than sporadic AD, in which other factors such as inflammation or immune response may be involved.

2.4.6 Specific genetic variants confer a distinctive cell composition profile

A variety of genetic variants increase risk of LOAD; however, it is unclear if the cellular mechanisms are the same across these distinct risk factors. Therefore, I tested the hypothesis that distinct genetic causes of LOAD have characteristic cellular population signatures.
I initially ascertained the effect of *APOE* ε4 on the cell-type composition. I observed a significant lower relative proportion of neurons (β = -0.06 for each of the ε4 alleles; p = 9.91×10^{-03}) and increase of relative proportion of astrocytes (β = 0.10; p = 4.15×10^{-02}) from the TC included in the Mayo Clinic dataset (*Table 2.20; Figure 2.28a; Figure 2.29a*). This finding was replicated when I performed a multi-region analysis of the MSBB dataset (β = -0.04; p = 2.60×10^{-03} and β = 0.05; p = 1.31×10^{-03} for neurons and astrocytes respectively; *Table 2.16; Figure 2.28a; Table 2.20; Figure 2.29a*). Given the strong risk conferred by the *APOE* ε4 allele[56], I studied its effects on the cell-type composition by restricting my analysis to AD brains. I observed a significant association in the multi-area analysis of the MSBB dataset (β = -0.03 p = 4.01×10^{-02}; *Table 2.16; Figure 2.28b; Table 2.21; Figure 2.29b*) and also a significant increase in relative proportion of astrocytes (β = 0.03; p = 1.23×10^{-02}; *Table 2.16; Figure 2.28b; Table 2.21; Figure 2.29b*). I also observed a significant decrease in relative proportion of neurons (β = -0.06; p = 2.11×10^{-02}; *Table 2.16; Figure 2.28c*) when I analyzed the LOAD and control brains from the Knight-ADRC. When I restricted the analysis to AD brains from the Knight-ADRC and compared the *APOE* ε4 carriers (N = 46) to non-carriers (N = 41) I also observed decreased relative neuronal proportion (β = -0.06; p = 2.69×10^{-02}; *Table 2.16; Figure 2.28d*). I extended the models to correct for the Braak stages, and observed a significant association for the relative proportion of neurons with the *APOE* ε4 allele in the Knight-ADRC dataset (β = -0.06; p = 3.66×10^{-02}; *Table 2.16*), and a significant association for the relative proportion of astrocytes in the MSBB (β = 0.04; p = 4.89×10^{-02}; *Table 2.16*). Furthermore, I performed a meta-analysis to combine the evidence of both studies and observed a significant association of the relative
neuronal proportion with APOE ε4 allele (p=1.86×10^{-02}) and marginally significant association for the relative astrocytic relative proportion (p=0.09).

Figure 2.28 Effect of the APOE ε4 allele and TREM2 coding variants on the cellular population structure. Mean neuronal (blue) and astrocytic (red) proportions for a) AD cases and controls in the Knight-ADRC brains categorized by APOE ε4 carriers vs. non-carriers and b) AD cases of Knight-ADRC brain bank (bars indicate standard deviations). c) AD cases and controls in the Mayo Clinic and MSBB d) AD cases in the Mayo Clinic and MSBB. e) Neuronal (blue) and astrocyte (red) distributions for samples included in the Mount Sinai brain bank stratified by TREM2 genetic status. APC: Anterior Prefrontal Cortex; STG: Superior Temporal Gyrus; PHG: Parahippocampal Gyrus; IFG: Inferior Frontal Gyrus; (n.s. P > 0.05; * P < 0.05; **** P < 1.0×10^{-4})
Figure 2.29 Neurons and astrocytes distributions for samples included in the Mayo Clinic and Mount Sinai Brain Bank (MSBB) stratified by APOE ε4 allele. Neuronal (blue) and astrocyte (red) proportions. a) AD Cases and controls. b) Restricted to AD cases.
Table 2.16 Gene specific cellular proportion analysis for Knight-ADRC and Mount Sinai Brain Bank studies

<table>
<thead>
<tr>
<th>Variant Carriers</th>
<th>Sample Size</th>
<th>Neuron</th>
<th>Effect</th>
<th>P-value</th>
<th>Astrocyte</th>
<th>Effect</th>
<th>P-value</th>
<th>Oligodendrocyte</th>
<th>Effect</th>
<th>P-value</th>
<th>Microglia</th>
<th>Effect</th>
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<tr>
<td>Knight-ADRC</td>
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<tr>
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<td>Sporadic AD vs Control</td>
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<tr>
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<td>673</td>
<td>0.04</td>
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<tr>
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<td>0.05</td>
<td>1.3×10^{-02}</td>
<td>-0.05</td>
<td>2.7×10^{-02}</td>
<td>-1.82×10^{-03}</td>
<td>8.13×10^{-01}</td>
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<td>1.28×10^{-01}</td>
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<tr>
<td>Mean plaque counts corrected</td>
<td>673</td>
<td>0.05</td>
<td>2×10^{-02}</td>
<td>-0.05</td>
<td>1.59×10^{-02}</td>
<td>1.73×10^{-03}</td>
<td>8.15×10^{-01}</td>
<td>-2.2×10^{-03}</td>
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<tr>
<td>APOE e4 counts all samples</td>
<td>556</td>
<td>-0.04</td>
<td>2.6×10^{-03}</td>
<td>0.05</td>
<td>1.31×10^{-03}</td>
<td>-0.01</td>
<td>4.47×10^{-02}</td>
<td>-3.58×10^{-04}</td>
<td>6.53×10^{-01}</td>
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<tr>
<td>APOE e4 counts AD cases</td>
<td>225</td>
<td>-0.03</td>
<td>4.01×10^{-02}</td>
<td>0.03</td>
<td>4.23×10^{-02}</td>
<td>-4.52×10^{-03}</td>
<td>3.73×10^{-01}</td>
<td>-5.13×10^{-04}</td>
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<td>CDR corrected</td>
<td>225</td>
<td>-0.03</td>
<td>2.02×10^{-02}</td>
<td>0.03</td>
<td>2.03×10^{-02}</td>
<td>-4.86×10^{-03}</td>
<td>3.19×10^{-01}</td>
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<tr>
<td>Braak corrected</td>
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<td>7.35×10^{-02}</td>
<td>0.04</td>
<td>4.89×10^{-02}</td>
<td>-0.01</td>
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<td>-1.08×10^{-03}</td>
<td>4.12×10^{-01}</td>
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</table>
Next, I analyzed the cellular composition in PLD3 carriers (N = 33). PLD3 carriers exhibited significantly lower relative proportion of neurons compared to controls (β = -0.10; p = 1.60×10^{-04}; Figure 2.27d) and a significant higher relative proportion of astrocytes (β = 0.13; p = 2.84×10^{-03}; Table 2.16; Figure 2.27d). Sporadic AD non-carrier cases also exhibited significantly lower relative proportion of neurons compared to controls (β = -0.11; p = 5.45×10^{-03}) and significant higher relative proportion of astrocytes (β = 0.13; p = 2.95×10^{-04}; Table 2.16; Figure 2.27d). The cell proportion between sporadic AD non-carriers and PLD3 carriers did not show any significantly difference (p > 0.05).

Finally, I performed similar analyses with TREM2 carriers. TREM2 is involved in the immune response and its role in amyloid-β deposition or clearance remain controversial[263]. My analysis on the Knight-ADRC data showed significantly higher relative astrocytic proportion in AD affected TREM2 carriers (N = 20) compared to controls (β = 0.11; p = 1.05×10^{-02}; Table 2.16; Figure 2.27d). Despite TREM2 carriers presented lower neuron relative proportion compared to controls, this difference was not statistically significant (p > 0.05; Table 2.16; Figure 2.27d). I analyzed whether the TREM2 carriers provided sufficient power to detect a significant association. My empirical estimates showed that TREM2 sample size provides 96% of power to detect an association with an effect size comparable to that observed for sporadic AD (β = -0.11). I also investigated the cellular proportion of the eleven TREM2 carriers in the MSBB dataset. The multi-region analysis showed TREM2 carriers do not show a significant difference in relative neuronal proportion compared to controls (p > 0.05; Table 2.16; Figure 2.28e), whereas in the AD TREM2 non-carriers the relative neuronal and astrocytic proportions are significantly different from controls (β = -0.07; p = 1.91×10^{-08} and β = 0.08; p = 1.25×10^{-08} respectively; Table 2.16; Figure 2.28e).
In fact, my analyses indicate that *TREM2* carriers have a unique cellular brain composition distinct than the other AD cases. *TREM2* brains showed significantly higher relative neuronal proportion ($\beta = 0.05; p = 1.98 \times 10^{-02}$) and significantly lower relative astrocyte proportion than the AD non-carries ($\beta = -0.05; p = 1.58 \times 10^{-02}$; Table 2.16). The distribution of CDR, mean number of amyloid plaques and Braak staging do not differ between strata. Nonetheless, I verified that the cellular proportions were still significant after correcting for each of those variables (Table 2.16). These results suggested that the mechanism that lead to disease in *TREM2* carriers is less neuron-centric than in the general AD population.

### 2.5 Discussion

I have developed, optimized and validated a digital deconvolution approach to infer cell composition from bulk brain gene expression that integrates publicly available cell-type specific expression data while addressing the heterogeneity of the phenotypic differences of samples and technical characteristics of transcriptome ascertainment. I acknowledge that the accuracy of this platform might be affected by the phenotypic diversity of the reference panel or the disease-induced dysregulation of genes it includes. However, the deconvolution approach proved to be robust to the genes included in the reference panel, as I demonstrated that the proportions it inferred are not driven by the expression of any single gene. This platform produced reliable cell proportion estimates, as was shown by the evaluation of independent datasets of iPSC-derived neurons and microglia, mice cortical neurons (Figure 2.22) and simulated chimeric libraries.

I used this approach to deconvolve studies that include large number of neuropathologically defined AD and control brains with their transcriptome ascertained in distinct brain regions and observed consistently significant lower relative neuronal proportion and increased relative astrocyte proportions in the cerebral cortex suggesting neuronal loss and
astrocytosis. Compatible with other studies, I also identified that the altered cellular proportion is also significantly associated with decline in cognition and Braak staging [239]. In contrast, I did not identify a significant difference in the cellular population structure in the cerebellum, a region not affected in AD (Table 2.10; Figure 2.25a).

We generated RNA-Seq data from brains carrying pathogenic mutations in APP, PSEN1, PSEN2, which cause alterations in Aβ processing and lead to ADAD, and also generated RNA-Seq from brains of LOAD and neuropath-free controls. I observed altered cell composition in both ADAD and LOAD compared to controls. However, I identified that ADAD brains have a different cell-type composition than disease-stage-matched LOAD, as the ADAD has a significantly lower relative neuronal proportion and more pronounced astrocytosis. Given the specific cellular population structure of the TREM2 carriers, I compared the neuronal and astrocytic relative proportion of ADAD to that of LOAD non-carriers of variants in TREM2 and observed significant differences (β = -0.09 and p = 6.89×10⁻³ for neurons and β = 0.10; p = 1.49×10⁻³ for astrocytes). This indicates that the difference of the relative proportion between ADAD and LOAD are not driven by TREM2 carrier brains. Based on my results, I would hypothesize that this change in Aβ processing of ADAD would lead to more direct to neuronal death than the pathological processes of LOAD. Similarly, decreased neuronal and increased astrocyte relative proportions were significantly associated with APOE ε4 allele. It has been reported APOE ε4 allele increase the risk for AD by affecting APP metabolism or Aβ clearance [44, 151], suggesting a direct link between APP metabolism and neuronal death.

In contrast, the analysis of the Knight-ADRC brains showed that the neuronal relative proportion decrease is less pronounced in TREM2 carriers than in other LOAD cases. I replicated this finding in a multi-area analysis from the MSBB dataset. These results may implicate that
TREM2 risk variants lead to a cascade of pathological events that differ from those occurring in sporadic AD cases, which is also consistent with the known biology of TREM2. Further longitudinal neuroimaging analysis are required to validate my findings. TREM2 is involved in AD pathology through microglia mediated pathways, implicated on altered immune response and inflammation[54]. Recent studies in TREM2 knock-out animals showed that fewer microglia cells were found surrounding Aβ plaques with impaired microgliosis[275]. Furthermore, TREM2 deficiency was reported to attenuate tauopathy against brain atrophy[168]. I found no significant difference in the proportion of microglia between AD cases and controls. However, I found significantly decreased microglia in brains exhibiting pathological aging (Table 2.12; Figure 2.24), proving that these studies are sufficiently powered to identify significant differences. In any case, I cannot rule out the possibility of a change in the activation stage of microglia in these individuals. Overall, these results suggest that TREM2 affects AD risk through a slightly different mechanism to that of ADAD or LOAD in general. Therefore, other pathogenic mechanisms should contribute to disease. I believe that a detailed modeling of immune response cells, reflecting the alternative microglia activation states, will generate more accurate profiles to elucidate the immune cell distribution in AD.

2.6 Conclusions

There is a large interest in the scientific community to use brain expression studies to try to identity novel pathogenic mechanism in AD and to identify novel therapeutic targets. These efforts are generating a large amount of bulk RNA-Seq data, as single-cell RNA (scRNA-Seq) from human brain tissue in large sample size is not feasible. Single-cell sorting needs to be performed with fresh tissue[115], which restrains the analysis of highly characterized fresh-frozen brains collected by AD research centers. My results indicate that digital deconvolution
methods can accurately infer relative cell distributions from brain bulk RNA-Seq data, but I recognize the importance of obtaining traditional neuropathological measures to validate the results I observed. Having this approach validated for AD can have an important impact in the community, because digital deconvolution analyses 1) can reveal distinct cellular composition patterns underlying different disease etiologies; 2) can provide additional insights about the overall pathologic mechanisms underlying different mutations carriers for variants as in genes such as TREM2, APOE, APP, PSEN1 and PSEN2; 3) can correct the effect that altered cell composition and genetic statuses have in addition to downstream transcriptomic analyses and lead to novel and informative results; 4) can help the analysis of highly informative frozen brains collected over the years.

In conclusion, my study provides a reliable approach to enhance our understanding of the fundamental cellular mechanisms involved in AD and enable the analysis of large bulk RNA-Seq data that may lead to novel discoveries and insights into neurodegeneration.
Chapter 3: The *TMEM106B* rs1990621 protective variant is associated with increased neuronal proportion
3.1 Abstract

**Background:** In previous studies, I observed decreased neuronal and increased astrocyte proportions in AD cases in parietal brain cortex by using a deconvolution method for bulk RNA-Seq. These findings suggested that genetic risk factors associated with AD etiology have a specific effect in the cellular composition of AD brains. The goal of this study is to investigate if there are genetic determinants for brain cell compositions.

**Methods:** Using cell type composition inferred from transcriptome as a disease status proxy, I performed cell type association analysis to identify novel loci related to cellular population changes in disease cohort. We imputed and merged genotyping data from seven studies in total of 1,669 samples and derived major CNS cell type proportions from cortical RNA-Seq data. I also inferred RNA transcript integrity number (TIN) to account for RNA quality variances. The model I performed in the analysis was: normalized neuronal proportion ~ SNP + Age + Gender + PC1 + PC2 + median TIN.

**Results:** A variant rs1990621 located in the TMEM106B gene region was significantly associated with neuronal proportion (p=6.40×10^{-07}) and replicated in an independent dataset. The association passed genome-wide multiple test correction in the multi-tissue meta-analysis (p=9.42×10^{-09}) and joint analysis (p=7.66×10^{-10}). This variant is in high LD with rs1990622 (r^2 = 0.98) which was previously identified as a protective variant for FTD with TDP-43 inclusion. Further analyses indicated that this variant is associated with increased neuronal proportion in participants with neurodegenerative disorders, not only in AD cohort but also in cognitive normal elderly cohort. However, this effect was not observed in a younger schizophrenia cohort with a mean age of death < 65. The second most significant loci for neuron proportion was
APOE, which suggested that using neuronal proportion as an informative endophenotype could help identify loci associated with neurodegeneration.

**Conclusion:** This result suggested a common pathway involving TMEM106B shared by aging groups in the present or absence of neurodegenerative pathology may contribute to cognitive preservation and neuronal protection.

3.2 Introduction

3.2.1 Alzheimer’s disease in the context of multi-cell type interactions

Although neuronal loss and synapse dysfunction are the preceding events of cognitive deficits in Alzheimer’s disease (AD), neurons do not work or survive by themselves. These delicate organelles require supports through intimate collaborations within themselves and with other cell types[125]. The microenvironment of cellular crosstalk, interaction, balance, and circuits maintained by neurons, astrocytes, microglia, oligodendrocytes, and other vascular cells are essential for the brain to carry out functions and fight against insults.

AD associated risk factors identified across the genome also point to the involvements of multi-cell types apart from neurons[125, 161]. APOE4 is related to lipid metabolism and mostly expressed in astrocyte and microglia[56]. Other lipid metabolism related risk genes are ABCA7 identified in all cell types[130, 161], CLU in astrocyte and oligodendrocyte precursor cells[119, 160, 161], and SORL1 in astrocyte[161]. Research interests in the roles of inflammatory response to toxic stimuli or microbial infection have been escalating recently, and AD risk genes associated with immune response including TREM2[114, 144, 246], PLCG2[246], ABI3[246], CR1[160, 161], CD33[130, 161], HLA-DRB5–HLA-DRBI[161], and INPP5D[161] are mostly expressed in microglia and macrophages. BIN1 expressed in microglia, oligodendrocyte, and
neurons[161], and *PICALM* expressed in microglia and endothelial cells[119, 161] are associated with endocytosis.

In a normal functional brain, astrocytes, microglia, and oligodendrocytes provide trophic supports to neurons and various cell type specific functions. Astrocytes confer multiple functions to fulfill neurons’ metabolic needs[250] including but not limited to providing substrates for oxidative phosphorylation[210], exerting regulation of excitatory CNS neurotransmitter glutamate[76, 93], and serving as bidirectional communication nodes that talk to both neurons and blood vessels and modulate their activities in an arrangement of functional entities named neurovascular units[227, 244, 260]. Microglia surveil in the extracellular space and look for pathogens or debris to engulf through phagocytosis. Oligodendrocyte provides insulation to neurons by wrapping around the axons with myelin sheath. However, in an AD diseased brain, these supporting cells may become double-edged swords that play beneficial and/or harmful roles as disease progresses. Amyloid-β accumulation and clearance are the central events of the amyloid cascade hypothesis. Both astrocyte and microglia have been involved in response to the toxic stimuli of amyloid plaques. During the early stage, microglia[124, 126, 127] and astrocytes[126, 209, 225] accumulate around plaques to phagocytose or degrade those in a protective manner. However, as disease progresses, the chronic and prolonged activation of microglia and astrocytes will be provoked into a damaging pro-inflammatory state and a vicious circle that exacerbate pathology in a harmful manner. Evidence suggested that increased inflammatory cytokine secretion in microglia, and increased production of complement cascade components, and impaired glutamate regulation (unregulated glutamate activity can cause neuronal excitatory cell death)[76] may contribute to synaptic loss which ultimately leads to cognitive deficits. Disrupted neuronal plasticity due to myelin loss and dysfunctional
neurovascular units further exacerbate the dreadful situation and destroy the harmony of the multi-cell type microenvironment.

3.2.2 Cell type composition inferred from bulk RNA-Seq deconvolution

Apart from disturbed homeostatic processes and impaired circuits integrity, cell type composition or proportion is also altered. Brains affected by AD exhibits neuronal loss, oligodendrocyte loss, astrocytosis, and microgliosis. However, the specific effects that pathological mutations and risk variants have on brain cellular composition are often ignored. To investigate the changes of cerebral cortex cell-type population structure and account for the associated confounding effects in downstream analysis, I developed an in-silico deconvolution method to infer cellular composition from RNA-Seq data, which has been documented in my previous publication[172], and explained in depth in Chapter 2. In summary, I firstly assembled a reference panel to model the transcriptomic signature of neurons, astrocytes, oligodendrocytes and microglia. The panel was created by analyzing expression data from purified cell lines. I evaluated various digital deconvolution methods and selected the best performing ones for my primary analyses. I tested the digital deconvolution accuracy on induced pluripotent stem cell (iPSC) derived neurons and microglia, and neurons derived from Translating Ribosome Affinity Purification followed by RNA-Seq. Finally, I verified its accuracy with simulated admixture with pre-defined cellular proportions.

Once the deconvolution approach was optimized, I calculated the cell proportion in AD cases and controls from different brain regions of LOAD and ADAD datasets. I found that neuronal and astrocyte relative proportions differ between healthy and diseased brains, and also differ among AD cases that carry different genetic risk variants. Brain carriers of pathogenic mutations in APP, PSEN1 or PSEN2 presented lower neuronal and higher astrocytes relative
proportions compared to sporadic AD. Similarly, APOE ε4 carriers also showed decreased neuronal and increased astrocyte relative proportions compared to AD non-carriers. In contrast, carriers of variants in TREM2 risk showed a lower degree of neuronal loss than matched AD cases in multiple independent studies. These findings suggest that different genetic risk factors associated with AD etiology may have gene specific effects in the cellular composition of AD brains.

3.2.3 Use cell type composition in cell type QTL (cQTL) to identify novel loci for AD risk

In a recently published study named PsychENCODE[272], a very similar deconvolution approach as reported in my previous study[172] was taken to infer cell type composition from RNA-Seq data predominantly drawn from psychiatric disorder cohorts. From the cell fractions inferred from bulk RNA-Seq data, they found that cell type composition differences can account for more than 88% of bulk tissue expression variation observed across the population with a ±4% variance on a per-subject level. Using cell type compositions as quantitative traits, the authors identified a non-coding variant closed to the FZD9 gene that is associated with both FZD9 gene expression and the proportion of excitatory layer 3 neurons[272]. Interestingly, deletion variants found previously upstream of FZD9 were associated with cell composition changes in Williams syndrome[45], a developmental disorder exhibits mild to moderate intellectual disabilities with learning deficits and cardiovascular problems. This observation re-emphasized the importance of incorporating cell type composition into RNA-Seq analysis pipeline even in psychiatric disorder cohorts without dramatic changes in cellular composition, not mention the necessity of such practice in neurodegeneration disorders that have significant changes in cell type composition. It also demonstrated the great potential of using relative abundance of specific cell types in
identifying novel variants and genes implicated in disease. However, it is unclear if this finding is only applicable to psychiatry-relate traits or it is a more general finding.

In this study, I utilized cell-type proportions inferred from my deconvolution method[172] to perform cell type QTL analysis in a dataset enriched for AD cases in search for potential new loci that are associated with neurodegeneration disorders. We imputed and merged genotyping or whole genome sequencing data from seven studies - five centered on neurodegeneration (N = 1,125), one schizophrenia cohort (N = 414), and GTEx multiple tissue controls (N = 130). From cortical RNA-Seq data, I derived cell fractions of four major CNS cell types, including neuron, astrocyte, microglia, and oligodendrocyte. Using normalized neuronal proportion as quantitative trait, I identified a variant rs1990621 located in the TMEM106B gene region significantly associated with neuronal proportion variation in all cohorts except schizophrenia subjects. This variant is in high LD with rs1990622 (r² = 0.98), which was previously identified as a protective variant in FTD cohorts[266]. Variants in this region have also been found to be associated with AD with TDP-43 pathology[229], and downregulation of TMEM106B is observed in AD brains[234] . In conclusion, I have identified a variant associated with neuronal proportion with potential protective effect in neurodegeneration disorders.

3.3 Methods

3.3.1 Study participants

The participants were sourced from seven studies with a total sample size of 1,669 (Table 3.1). Among those, five studies are mainly focused on neurodegenerative disorders including Alzheimer’s disease (N = 681), frontotemporal dementia (N = 11), progressive supranuclear palsy (N = 82), pathological aging (N = 29), Parkinson Disease (N = 1), as well as cognitive normal individuals (N = 540). These samples come from the Mayo, MSSM, Knight
ADRC, DIAN, and ROSMAP studies as described in table 3.1. To compare with the neurodegenerative disorders, I also included schizophrenia (N = 210) and bipolar disorders (N = 34) participants from the CommonMind study (Table 3.1). Additionally, two studies, MSSM and GTEx, contain multi-tissue data that include some participants contribute more than one tissue (Table 3.1).

3.3.2 Standard protocol approvals, registrations and patient consents

The protocol of DIAN and Knight-ADRC studies have been approved by the review board of Washington University in St. Louis. The protocol of Mayo dataset was approved by the Mayo Clinic Institutional Review Board (IRB). All neuropsychological, diagnostic and autopsy protocols of MSSM dataset were approved by the Mount Sinai and JJ Peters VA Medical Center Institutional Review Boards. The religious orders study and the memory and aging project of ROSMAP was approved by the IRB of Rush University Medical Center. The NIH Common Fund’s GTEx program protocol was reviewed by Chesapeake Research Review Inc., Roswell Park Cancer Institute’s Office of Research Subject Protection, and the institutional review board of the University of Pennsylvania. Within CommonMind consortium, the MSSM sample protocol was approved by Icahn School of Medicine at Mount Sinai IRB; the Pitt sample protocol was approved by the University of Pittsburgh’s Committee for the Oversight of Research involving the Dead and IRB for Biomedical Research; the Penn sample protocol was approved by the Committee on Studies Involving Human Beings of the University of Pennsylvania. All participants were recruited with informed consent for research use.
Table 3.1 Demographic information for cohorts included in the study. AD: Alzheimer’s Disease; FTD: frontal temporal dementia; PSP: progressive supranuclear palsy; PA: pathological aging; PD: Parkinson’s Disease; SCZ: schizophrenia; BP: bipolar disease; OTH: other unknown dementia or no diagnosis information.

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<th>FTD</th>
<th>PSP</th>
<th>PA</th>
<th>PD</th>
<th>SCZ</th>
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Table 3.2 General information of seven studies evolved in the analysis. TCX: temporal cortex; PAR: parietal cortex; CTX: cortex; FCX: frontal cortex; DLPFC: dorsal lateral prefrontal cortex. BM9: dorsal lateral prefrontal cortex; BM10: Anterior prefrontal cortex; BM22: superior temporal gyrus; BM24: ventral anterior cingulate cortex; BM36: parahippocampal gyrus; BM44: inferior frontal gyrus. Mean coverage unit is million.

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<td>Single end</td>
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<td>RNA depletion</td>
<td>HiSeq 2500</td>
<td>35.96 ± 10.04</td>
<td>WGS</td>
<td>Wang 2018</td>
</tr>
<tr>
<td>Knight-ADRC</td>
<td>PAR</td>
<td>Paired end</td>
<td>150</td>
<td>RNA depletion</td>
<td>HiSeq 4000</td>
<td>137.87 ± 21.81</td>
<td>Genotype</td>
<td>Li 2018</td>
</tr>
<tr>
<td>DIAN</td>
<td>PAR</td>
<td>Paired end</td>
<td>150</td>
<td>RNA depletion</td>
<td>HiSeq 4000</td>
<td>149.82 ± 19.68</td>
<td>Genotype</td>
<td>Li 2018</td>
</tr>
<tr>
<td>GTEx</td>
<td>BM24 CTX FCX</td>
<td>Paired end</td>
<td>76</td>
<td>ploy-A selection</td>
<td>HiSeq 2000</td>
<td>48.28 ± 13.2</td>
<td>WGS</td>
<td>GTEX 2013; Battle 2017</td>
</tr>
<tr>
<td>CommonMind</td>
<td>BM9</td>
<td>Paired end</td>
<td>100</td>
<td>RNA depletion</td>
<td>HiSeq 2500</td>
<td>86 ± 21.12</td>
<td>Genotype</td>
<td>Fromer 2016</td>
</tr>
</tbody>
</table>


113
3.3.3 Data collection and generation

Cortical tissues from various locations of post-mortal brains were collected (Table 3.2). RNA was extracted from lysed tissues and prepared into libraries of template molecules ready for subsequent next-generation sequencing steps. Ribosomal RNAs constitute 80%-90% of total RNAs, which are not the targets of this study. To focus on mRNA quantification usually researchers would either remove excessive rRNAs or enrich for mRNAs during RNA-Seq library preparation. In this study, DIAN[172], Knight ADRC[172], MSSM[274], and CommonMind[95] took a rRNA depletion approach to removed ribosomal RNA from total RNAs to retain a higher mRNA content. Whereas, Mayo[9], ROSMAP[27, 28], and GTEx[6, 24] took a poly-A enrichment approach to enrich mRNAs from total RNAs. Genotype information were also collected and sequenced correspondingly. RNA-Seq paired with genotype data for each participant were either sequenced at Washington University for DIAN and Knight-ADRC studies or downloaded from public database for all the other studies. Please see Table 3.2 and each study reference(s) for more data collection and generation specifications.

3.3.4 Data QC and preprocessing

Genetic Data

Stringent quality control (QC) steps were applied to each genotyping array or sequence data. The minimum call rate for single nucleotide polymorphisms (SNPs) and individuals was 98% and autosomal SNPs not in Hardy-Weinberg equilibrium (p-value < 1x10^-6) were excluded. X-chromosome SNPs were analyzed to verify gender identification. Unanticipated duplicates and cryptic relatedness (Pihat ≥ 0.25) among samples were tested by pairwise genome-wide estimates of proportion identity-by-descent. EIGENSTRAT[215] was used to calculate principal components. The 1000 Genomes Project Phase 3 data (October 2014),
SHAPEIT v2.r837[67], and IMPUTE2 v2.3.2[134] were used for phasing and imputation. Individual genotypes imputed with probability < 0.90 were set to missing and imputed genotypes with probability ≥0.90 were analyzed as fully observed. Genotyped and imputed variants with MAF < 0.02 or IMPUTE2 information score < 0.30 were excluded. WGS data quality is censored by filtering out reads with sequencing depth DP < 6 and quality GQ < 20 followed by similar QC approaches as described above for genotyping data. After the QC, all studies including imputed genotype and WGS data was merged into a binary file using Plink for downstream analysis. PCA and IBD analyses were performed on the merged binary files using Plink to keep European ancestry and unrelated participants (Figure 3.1 and Figure 3.2).

**Figure 3.1 Genomic PCA analysis.** Genotype data PCA analysis was performed to select European ancestry subjects with PC1 < -0.002 and PC2 < 0.008 with red dotted cut-off lines. HapMap_CEU: HapMap Utah residents with Northern and Western European ancestry; HapMap_JPT: HapMap Japanese in Tokyo, Japan; HapMap_YRI: HapMap Yoruba in Ibadan, Nigeria; MayoADGS: Mayo Clinic study participants; MSBB: MSSM study participants; GTEx: GTEx study participants; DIAN: DIAN study participants; MAP: Knight-ADRC participants; NIALOAD: Knight-ADRC participants; CMC: CommonMind participants; ROSMAP: ROSMAP participants.
Expression Data

FastQC was applied to RNA-Seq data to examine various aspects of sequencing quality[231]. Outlier samples with high rRNA contents or low sequencing depth were removed from the pool. The remaining samples were aligned to human GRCh37 primary assembly using Star with 2-Pass Basic mode (ver 2.5.4b)[74]. Alignment metrics were ascertained by applying Picard CollectRnaSeqMetrics[4] including reads bias, coverage, ribosomal contents, coding bases, and etc. Following which, transcript integrity number (TIN) for each transcript was calculated on aligned bam files using RSeQC tin.py[273] (ver 2.6.5). RNA-Seq coding gene and transcript expression was quantified using Salmon transcript expression quantification (ver 0.7.2) with GENCODE Homo sapiens GRCh37.75 reference genome[208].

Figure 3.2 Genomic IBD analysis. IBD analysis was performed to select unrelated subjects with $Z_0 > 0.8$ and $Z_1 < 0.2$ with red dotted cut-off lines. When there are related individuals, one individual will be dropped from the related pair.
Four major central nerve system cell type proportions were inferred from RNA-Seq gene expression quantification output as documented in my previous deconvolution study[172]. To briefly explain the deconvolution process, I firstly assembled a reference panel to model the transcriptomic signature of neurons, astrocytes, oligodendrocytes and microglia from purified single cell tissue sources respectively. Using the reference panel and the method population-specific expression analysis[157] (PSEA, also named meanProfile in CellMix implementation[102]), I calculated four cell type proportions for each subject bulk RNA-Seq data. For each brain tissue collection site of each study, outlier values for each cell type proportion were removed. Mean values for each cell type of each tissue in each study were subtracted from the deconvolution results to center all the distributions to zero mean (Figure 3.3). Phenotype information from all studies were merged and unified to the same coding paradigm to enable downstream joint analysis; for example, males are all coded as 1 and females are 2.
Figure 3.3 Major CNS cell type proportions derived from RNA-Seq datasets with each row representing each tissue of each study. A) raw cell type proportions inferred from the data with vertical bars indicating quantiles within each tissue and each cell type. B) cell type proportions were normalized by subtracting the mean from each tissue deconvolution result after removing outliers.
3.3.5 Data analysis

For the discovery phase, ROSMAP dataset was analyzed with linear regression model employed in Plink\cite{217} using normalized neuronal proportion to run quantitative trait analysis. Age, sex, PC1, PC2, and median TIN were used as covariates to account for potential genetic, phenotypic or technical heterogeneity. TIN is calculated directly from post-sequencing results that captures RNA degradation by measuring mRNA integrity directly\cite{273}. Results were depicted as Manhattan plots using R (ver 3.4.3) qqman package\cite{261} (ver 0.1.4).

For the replication phase, all the other studies except ROSMAP were combined and prepressed to run meta-tissue QTL analysis because MSSM and GTEx contain samples with multiple cortical tissues. Meta-Tissue software installation and data preprocessing were conducted following a four-step instruction documented in the developer website: [http://genetics.cs.ucla.edu/metatissue/install.html](http://genetics.cs.ucla.edu/metatissue/install.html). Meta-tissue\cite{253} processing pipeline calls two main functions, firstly MetaTissueMM\cite{253} and then followed by Metasoft\cite{117}. MetaTissueMM applies a mixed model to account for the heterogeneity of multiple tissue QTL effects. Metasoft performs the meta-analysis while proving a more accurate random effect p-value for multiple tissue analysis and a m-value based on Bayesian inference to indicate how likely a locus is a QTL in each tissue. Similarly, results were depicted as Manhattan plots and visually examined.

For the final merging phase, both discovery and replication studies were combined to maximize sample size. Apart from meta-tissue analysis by each tissue of each study, a split by disease status analysis was also performed in the final merging phase. Samples from each tissue of each study were also split into disease categories. Resultant subcategory with less than 20
subjects were removed from the analysis to avoid false results due to too small sample size. Similar data preparation and analysis pipeline were enforced as documented above.

QTL analysis results were uploaded to Fuma (v1.3.3d)[276] to annotation significant SNPs (p-value < 10^{-06}) with GWAScatalog (e91_r2018-02-06) and ANNOVAR (updated 2017-07-17). Gene-based analysis was also performed by Magma (v1.06)[63] implemented in Fuma.

3.3.6 Data availability

Mayo: https://www.synapse.org/#!Synapse:syn5550404

MSSM: https://www.synapse.org/#!Synapse:syn3157743

ROSMAP: https://www.synapse.org/#!Synapse:syn3219045

CommonMind: https://www.synapse.org/#!Synapse:syn2759792


Knight-ADRC: https://www.synapse.org/#!Synapse:syn12181323

According to the data request terms, DIAN data are available upon request: http://dian.wustl.edu

3.4 Results

3.4.1 Study design

The ROSMAP study containing 523 subjects will be the discovery dataset, and the other six studies are collapsed into replication dataset with 1,146 subjects. Altogether, I have assembled a set of cortical RNA-Seq data comprised of 1,669 participants predominantly focused on neurodegenerative disorders from seven sources (Figure 3.4, Table 3.1). Collectively, Mayo, MSSM, Knight ADRC, and ROSMAP studies contributed 664 sporadic AD cases. Apart from sporadic AD, 15 subjects from DIAN study and 2 from Knight-ADRC also
harbor *PSEN1*, *PSEN2*, and *APP* mutations that exhibit familial AD inheritance pattern. Other neurodegenerative disorders, including progressive supranuclear palsy (PSP), pathological aging (PA), frontal temporal dementia (FTD), and Parkinson’s Disease (PD), are mainly drawn from Mayo and Knight ADRC datasets. Other psychiatric disorders including schizophrenia and bipolar disorders are contributed by the CommonMind study. Besides, 540 control subjects cleared of cognitive dementia or neuropsychiatric symptoms were also included. MSSM and GTEx also included multiple tissue data, which were collected from multiple regions of the same subjects that allow us to perform region specific comparison within the same cohort.

Discovery analysis was performed in ROSMAP study. In the replication phase, all the other studies were merged to replicate signals identified from the discovery ROSMAP set. Because GTEx and MSSM contain multiple cortical regions collected from the same subjects, I also applied meta-tissue software[253] specifically designed for multi-tissue QTL analysis to perform a mixed model analysis with random effects that account for correlated measurements from multi-tissue individuals. To attain the largest available sample size for this study, the discovery and replication sets were merged to perform the merged multi-tissue QTL analysis in a search for additional signals hidden in previously separated discovery or replication analysis due to lack of power. After merged analysis, the cohorts were split into four major disease status groups (AD, control, schizophrenia, other non-AD neurodegenerative disorders) to explore how different disease strata could impact the results.
Figure 3.4 Study design. RNA-Seq and paired genotype or WGS data were accessed and preprocessed for downstream analysis. Genotype data was censored based on my quality control criteria and imputed as needed. WGS and imputed genotype were merged and followed by PCA and IBD procedures to select unrelated European ancestry subjects. RNA-Seq data was quality checked with FastQC and aligned to human GRCh37 primary assembly with Star, from which TIN was inferred with RSeQC to account for RNA integrity variances that I later incorporated into the analysis. Gene expression were quantified from unaligned RNA-Seq with pseudo-aligner Salmon for deconvolution procedure. Cell type composition comprised of four major CNS cell type proportions were inferred by performing deconvolution procedure on gene expression quantification results. Using cell type proportions as quantitative traits, I identified loci in TMEM106B gene region associated with neuronal proportion in my assembled dataset.
3.4.2 *TMEM106B* variants associated with neuronal proportion

During discovery phase, ROSMAP dataset (N = 484 after removing outliers from total number of 523 subjects) was used to perform cell type proportion QTL analysis. Using normalized neuronal proportion as a quantitative trait, the QTL analysis identified more than 10 peaks that passed genome wide suggestive threshold (<1.0×10⁻⁰⁵, Figure 3.5AB, Table 3.3). However, only one signal rs1990621 (chr7: 12283873) were replicated with p-value = 7.41×10⁻⁰⁴ in the replication dataset (N = 1,052) combining all the other datasets except ROSMAP (Figure 3.5CD). When the discovery and replication datasets were merged to attained a larger sample size (N = 1,536), rs1990621 major allele C is negatively associated with neuronal proportion with p-value = 9.42×10⁻⁰⁹ (Figure 3.6AB, Figure 3.7AC), which means the minor allele G is associated with increased neuronal proportion in my assembled datasets focusing on neurodegenerative disorders.

Table 3.3 rs1990621 (chr7:12283873) major allele C is significantly associated with decreased neuronal proportions. Therefore, G allele (MAF = 0.4658) is significantly associated with increased neuronal proportions.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Brain Region</th>
<th>Ref Allele</th>
<th>Sample Size</th>
<th>Beta</th>
<th>SE</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Discovery</td>
<td>DLPFC</td>
<td>C</td>
<td>484</td>
<td>-0.3</td>
<td>0.06</td>
<td>6.40×10⁻⁰⁷</td>
</tr>
<tr>
<td>Replication</td>
<td>Multiple</td>
<td>C</td>
<td>1,052</td>
<td>-0.13</td>
<td>0.04</td>
<td>7.41×10⁻⁰⁴</td>
</tr>
<tr>
<td>Merged meta-tissue</td>
<td>Multiple</td>
<td>C</td>
<td>1,536</td>
<td>-0.16</td>
<td>0.05</td>
<td>9.42×10⁻⁰⁹</td>
</tr>
</tbody>
</table>
Figure 3.5 Discovery and replication phases Manhattan and QQ plots. Loci located in chromosome 7 were associated with neuronal proportion in ROSMAP discovery dataset and replicated in replication dataset. A) Discovery set Manhattan plot showed seven peaks associated with neuronal proportion at suggestive threshold. The peak located in chromosome 7 was labeled, which is for rs1990621 with $p$-value $= 6.4 \times 10^{-07}$. B) QQ plot of the discovery phase analysis. C) Replication set Manhattan plot showed that the peak located in chromosome 7 replicated the signal identified during discovery phase with $p$-value $= 7.41 \times 10^{-04}$. D) QQ plot of the replication phase analysis.
**Figure 3.6 SNP-based and gene-based meta-analysis.** rs1990621 located in chromosome 7 TMEM106B gene region was significantly associated with neuronal proportion in cortical RNA-Seq dataset. A) Manhattan plot showed SNP-based genome-wide significant hit located in chromosome 7 with other suggestive SNP hits labeled. B) QQ plot of the SNP-based analysis. C) Manhattan plot showed gene-based genome-wide significant hit located in chromosome 7 with other suggestive gene hits labeled. D) QQ plot of the gene-based analysis.
Figure 3.7 Meta-Tissue analysis results of rs1990621. A) Forest plot showed p-value and confidence interval for rs1990621 for each tissue site of each dataset that included in the Meta-Tissue analysis. Summary random effect was depicted at the bottom as RE Summary. B) PM-Plot of rs1990621 while combining both p-value (y axis) and m-value (x axis). Red dot indicates that the variant is predicted to have an effect in that particular dataset, blue dot means that the variant is predicted to not have an effect, and green dot represents ambiguous prediction. C) Forest plot p-value and confidence interval for rs1990621 for discovery, replication, and merged datasets. D) Forest plot p-value and confidence interval for rs1990621 when splitting the merged dataset into four main disease categories.
Noticeably, in both replication and merged analyses, multi-tissue data were involved that provided additional power but also posed challenges to the analysis, the same issue faced by the GTEx study[24, 55]. Compared to a tissue-by-tissue approach, multiple tissues collected from the same subject may help identify QTL by aggregating evidence from multiple tissues, which is similar to a meta-analysis of combining each study. However, one violation of such approach is that the tissues collected from the same subject are presumably highly correlated since they shared the same genetic architecture. Thus, it violates the assumption of independency for carrying out a standard meta-analysis[253]. Another challenge of the multi-tissue QTL is the heterogeneity of the effects, which means a variant may have different effects on different tissues. To resolve these issues, I applied the Meta-Tissue analytic pipeline[253] (http://genetics.cs.ucla.edu/metatissue/) specifically designed for multi-tissue QTL, the same approach that GTEx took to analyze their multi-tissue data. As shown in Figure 3.7A, Meta-Tissue analysis results of rs1990621 for the merged analysis were displayed as a forest plot with 95% confidence interval and p-value labeled for each tissue of each study. Among them, MSSM and GTEx are multi-tissue studies while the others are single-tissue studies. Meta-Tissue used a linear mixed model to capture the multi-tissue correlation within MSSM and GTEx respectively. Regarding the effect heterogeneity, Meta-Tissue calculated a m-value[117] to predict if a variant has an effect in a tissue. M-value is similar to the posterior probability of association based on the Bayes factor[117] but with differences specifically designed for detecting whether an effect is present in a study included in a meta-analysis. Figure 3.7B is a PM-Plot that integrates evidences from both frequentist (p-value) and Bayesian (m-value) sides to interpret the heterogeneity of multi-tissue QTL effects. Variant rs1990621 in ROSMAP and Mayo studies have m-values greater than 0.9, are predicted to have an effect and color coded with red. In CMC
study, the m-value is less than 0.1, so it is predicted to not to have an effect and color coded with blue. All the other studies with m-value between 0.1 and 0.9 are predicted with ambiguous effect and color coded with green. Based on the forest plot and PM-Plot, the variant does have effect heterogeneity across different tissues and studies. In this case, random-effect model will be more suitable to account for effect heterogeneity. Therefore, summary random effect and p-value were reported for the analysis.

Apart from multi-tissue QTL, a single-tissue joint analysis was also performed. In this case, one tissue region was drawn from the multi-tissue data to avoid violating the independency assumption. Specifically, BM36 and frontal cortex tissue were selected to represent MSSM and GTEx study respectively. Study sites were coded as dummy variables to account for potential batch effects. In this joint analysis, the variant rs1990621 is also the top hit with p-value = 7.66×10^{-10}.

### 3.4.3 Neuronal protective effect of *TMEM106B* variants observed in neurodegenerative disorders and normal aging participants

To explore the effect in different disease categories, the merged dataset was stratified based on disease: AD, other non-AD neurodegenerative disorders, schizophrenia and control. Signification associations between rs1990621 and neuronal proportion were observed in AD (p-value = 1.95×10^{-07}), other non-AD neurodegenerative (p-value = 8.19×10^{-04}), and cognitive normal control (p-value = 2.94×10^{-02}) cohorts, but not in schizophrenic cohort (p-value = 9.32×10^{-01}, Table 3.4, Figure 3.7D). The effect of the variant was more prominent in neurodegenerative cohorts and aging controls with mean age of death greater than 65 years old. However, it was absent from younger cohorts such as GTEx controls and CommonMind
schizophrenia participants. Thus, this variant seems to be associated with a neuronal protection mechanism shared by any aging process in the present or absence of neuropathology.

### 3.4.4 Functional annotation of rs1990621

The variant rs1990621 is located in the *TMEM106B* gene region where other variants in high LD linkage are also located and labeled in Figure 3.8A. Although the CADD score and RegulomeDB score for this variant are not remarkably high to suggest any functional consequences (Figure 3.8BC), this variant is in high LD with rs1990622 ($r^2 = 0.98$), a *TMEM106B* variant previous identified to be associated with FTD risk[266], particularly in granulin precursor (*GRN*) mutation carriers[57, 92]. *TMEM106B* is expressed in neurons and microglia, with highest protein expression detected in the late endosome/lysosome compartments of neurons[36, 163, 237, 248]. A nonsynonymous variant rs3173615, which is also in high LD with rs1990621 ($r^2 = 0.98$), located in the exon 6 of *TMEM106B* (the dark blue dot in Figure 3.8B) produces two protein isoforms (p.T185S) that affect TMEM106B protein level through protein degradation mechanism[36, 49, 200].

<table>
<thead>
<tr>
<th>Disease</th>
<th>Brain Region</th>
<th>Ref Allele</th>
<th>Sample Size</th>
<th>Beta</th>
<th>SE</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
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<td>639</td>
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<td>0.07</td>
<td>$1.95\times 10^{-07}$</td>
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<tr>
<td>Control</td>
<td>Multiple</td>
<td>C</td>
<td>476</td>
<td>-0.14</td>
<td>0.06</td>
<td>$2.49\times 10^{-02}$</td>
</tr>
<tr>
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<td>BM9</td>
<td>C</td>
<td>189</td>
<td>-0.01</td>
<td>0.09</td>
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</tr>
<tr>
<td>Other</td>
<td>TCX</td>
<td>C</td>
<td>103</td>
<td>-0.45</td>
<td>0.14</td>
<td>$8.19\times 10^{-04}$</td>
</tr>
</tbody>
</table>

**Table 3.4** rs1990621 (chr7:12283873) major allele C is significantly associated with decreased neuronal proportions in AD, Control, and other non-AD neurodegenerative disorders. SCZ: schizophrenia; other: other non-AD neurodegenerative disorders, including progressive supranuclear palsy and pathological aging. BM9: dorsal lateral prefrontal cortex. TCX: temporal cortex.
Figure 3.8 Variant rs1990621 functional annotation and local plot. A) Local plot showed the zoom-in view of the hit in chromosome 7 with the top lead SNP rs1990621 labeled with dark purple. Nearby SNPs were also mainly located in the TMEM106B gene region and color coded with LD r² thresholds. B) Bottom panel showed combined CADD score, RegulomeDB score, and Chromatin state of the region shown in the top panel. C) Regulome DB and chromatin state explanation.
3.4.5 The impact of other neurodegenerative risk loci on neuronal proportion

To investigate what other AD or FTD variants might have an effect in neuronal proportion QTL analysis, I extracted results for 38 SNPs examined in two large scale genome wide association studies, AD focused (Lambert et al.[161]) and FTD focused (Ferrari et al.[87]) studies. Among those, only variants located in \textit{TMEM106B} and \textit{APOE} gene regions passed genome wide significant or suggestive threshold. Both rs1990622 (Figure 3.9A) and rs2075650 (Figure 3.9B) were found to be associated with FTD reported in Ferrari et al., which were associated with neuronal proportion in this study (Table 3.5). The top signals in \textit{APOE} region are rs283815, rs769449, and rs429358 with p-value < 1.22×10^{-05}. Note that rs429358 is one of the two SNPs that determine \textit{APOE} isoforms. Remember that \textit{APOE} \varepsilon4 alleles, coded by rs429358(C) and rs7412(C), confers the largest effect for AD risk. I observed that the C allele of rs429358 was associated with decreased neuronal proportion, but no association observed between rs7412 and neuronal proportion.

In a gene-based analysis of my neuronal proportion QTL, \textit{TMEM106B} (p-value = 2.96×10^{-08}) is the only gene that passed genome-wide significant threshold followed by \textit{APOE} (p-value = 3.2×10^{-05}), the most important gene for sporadic AD risk (Figure 3.6CD). Previous GWAS for AD risk performed with the International Genomics of Alzheimer’s Project (IGAP) data stratified by \textit{APOE} genotype showed that AD risk is significantly influenced by the interaction between \textit{APOE} and \textit{TMEM106B}[146]. Together with my observation of cellular composition QTL, these results suggest a potential interaction of \textit{TMEM106B} and \textit{APOE} may play a role in affecting AD risk/vulnerability and cellular composition balance between neurons and astrocytes, and the endosome and lysosome compartments might be the location that the interaction takes place.
<table>
<thead>
<tr>
<th>SNP</th>
<th>CHR</th>
<th>BP</th>
<th>Gene</th>
<th>Minor</th>
<th>Major</th>
<th>MAF</th>
<th>SNP proxy</th>
<th>cQTL Effect (Major)</th>
<th>AD risk p-value</th>
<th>OR (Minor)</th>
<th>FTD risk p-value</th>
<th>OR (Minor)</th>
<th>FTD-TDP risk p-value</th>
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<td>A</td>
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<td>1.18</td>
<td>5.7x10⁻²⁴</td>
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<td>-</td>
</tr>
<tr>
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<td>BTNL2</td>
<td>A</td>
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<td>-</td>
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<td>7.83x10⁻⁰⁴</td>
<td>-</td>
<td>-</td>
<td>0.775</td>
<td>1.57x10⁻⁰⁶</td>
</tr>
<tr>
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<td>A</td>
<td>C</td>
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<td>8.1x10⁻⁰⁴</td>
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<td>5.51x10⁻⁰⁵</td>
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<td>0.93</td>
<td>3.2x10⁻⁰⁸</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

Table 3.5 Neuronal proportion cQTL p-values were reported for variants previously identified in AD risk (by Lambert et al.), FTD risk (by Ferrari et al.), and FTD-TDP risk (by Van Deerlin et al.) studies.
Figure 3.9 **TMEM106B and TOMM40/APOE regions local plot.** A) Local plot showed the zoom-in view of the hit in chromosome 7 with target SNP rs1990622 labeled with dark purple, and the top leading SNP is rs1990621. Nearby SNPs were also mainly located in the **TMEM106B** gene region and color coded with LD r² thresholds. B) Local plot showed the zoom-in view of the hit in chromosome 19 with target SNP rs2075650 labeled with dark purple, and the top three leading SNPs are rs283815, rs769449, and rs429358. Nearby SNPs were also mainly located in the **TOMM40/APOE** gene region and color coded with LD r² thresholds. One gene omitted in this region is SNRPD2.
3.5 Discussions

The common variant rs1990622 in TMEM106B was first identified to be associated with FTD with TDP-43 inclusions[266]. Hyper-phosphorylated and ubiquitinated TDP-43 is the major pathological protein for FTD and ALS[198], which is also present in a broader range of neurodegenerative disorders, including AD[10], Lewy body disease[196], and hippocampal sclerosis[10]. Recent study also suggested distinct TDP-43 types present in non-FTD brains, typical TDP-43 α-type and newly characterized β-type[145]. TDP-43 α-type is the typical form conventionally observed in temporal, frontal and brainstem regions. TDP-43 β-type is characterized by its close adjacency to neurofibrillary tangles, which is predominantly observed in limbic system, including amygdala, entorhinal cortex, and subiculum of the hippocampus. These findings suggested that pathologic TDP-43 protein that closely associated with TMEM106B variants might be the common pathologic substrate linking these neurodegenerative disorders. Multiple lines of evidence have merged and shown that protective variants in TMEM106B are associated with attenuated cognitive deficits or better cognitive performance in ALS[268], hippocampal sclerosis[191], presymptomatic FTD[214], and aging groups with various neuropathological burden[280] or in the absence of known brain disease[222]. My study identified a protective variant rs1990621 of TMEM106B is associated with increased neuronal proportion in participants with neurodegenerative disorders and normal aging in non-demented controls. However, this effect is not observed in a younger schizophrenia cohort with a mean age of death less than 65 years old. This result suggested a common pathway involving TMEM106B shared by aging groups in the present or absence of neurodegenerative pathology that may contribute to cognitive preservation and neuronal protection.
My study has demonstrated that a protective variant rs1990621 identified in *TMEM106B* gene region may exert neuronal protection function in aging groups. A protein coding variant rs3173615 in high LD with rs1990621 ($r^2 = 0.98$) produces two protein isoforms (p.T185S). The S185 allele is protective and the protein carrying this amino acid is degraded faster than the risk variant T185. Thus, the risk allele of this coding variant leads to increased TMEM106B protein level[36, 49, 200]. *TMEM106B* overexpression results in enlarged lysosomes and lysosomal dysfunction[36, 295]. It has also been shown that TMEM106B may interact with PGRN (the precursor protein for granulin) in lysosome[200]. Although rs3173615 is not included in my genomic data, it is in complete linkage disequilibrium with rs1990621 and rs1990622. It is worth pointing out that the minor allele of rs1990622, which has a protective effect in FTD, is in-phase with the minor allele of rs1990621, which is associated with increased neuronal proportion in my analysis. Despite the fact that my dataset is focused on neurodegeneration, I only have 11 verified FTD cases suggesting that TMEM106B might have a general neuronal protection role in neurodegeneration apart from FTD.

This observation suggested that a potential involvement of *TMEM106B* in the endosome/lysosome pathway may play a role in neurodegenerative disorder risk or vulnerability. Neuronal survival requires continuous lysosomal turnover of cellular contents through endocytosis and autophagy[202]. Impaired lysosomal function reduces lysosomal degradative efficiency, which leads to abnormal build-up of toxic components in the cell. Impaired lysosomal system has been found to be associated with a broad range of neurodegenerative disorders, including AD[201], Parkinson disease[12, 233, 278], Huntington disease[90, 271], FTD[167], ALS[90], Niemann-Pick disease type C[154, 204], Creutzfeldt-Jakob disease[166], Charcot-Marie Tooth disease type 2B[243], Neuronal ceroid lipofuscinoses (Batten disease)[155, 156],
autosomal dominant hereditary spastic paraplegia[220], Chediak-Higashi syndrome[159], inclusion body myositis[14], and osteopetrosis[141]. Considering the extensive involvements of lysosomal/endosomal compartments in neurodegenerative disorders, it has been proposed that a long and chronic process of abnormal metabolic changes during aging has led to the accumulation of toxic materials[202]. When lifespan increases especially in the sporadic forms of neurodegenerative disorders, failures to degrade these waste products break the proteostasis and the balance maintained by the multicellular interactions, and trigger subsequent chain reactions that lead to neuronal death and outbreaks of various neurodegenerative disorders due to different genetic susceptibilities and other disease etiologies. Although each neurodegenerative disorder has its own characteristic proteopathy, the boundaries of protein pathology distribution are never clear-cut across different disorders. In fact, copathology or nonspecific pathology of proteopathy have been observed in most autopsies of neurodegenerative disorders, such as TDP-43 discussed above, Lewy body, α-synuclein[82], and etc. My observation of lysosomal gene TMEM106B associated with neuronal proportion in aging cohorts suggests that the lysosomal pathway might be involved in the common mechanism underlying a broad range of neurodegenerative disorders or aging process in general that contribute to neuronal cell death.

My study has demonstrated the great potential of using cell type composition as quantitative traits to identify QTLs associated with the changes in cell fractions. This approach is more powerful for disorders that involve considerably changes in cellular composition, for example, neurodegenerative disorders, and normal conditions during developmental or aging processes. The development of recent single cell studies will greatly increase the resolution in advancing our knowledge of cellular population changes. More detailed fine mapping of cellular composition from single cell studies together with machine learning algorithms, bulk RNA-Seq
deconvolution will be more accurately capturing cellular fraction changes in the samples, such as different types of neurons or different states of astrocytes or microglia. Regarding scalability, this single cell powered bulk deconvolution approach is preferable for carrying out such cell type composition QTL analysis, because due to the high cost of performing single cell studies, bulk RNA-Seq is more financially feasible to scale up, and with larger sample sizes more hidden signals will be unrevealed with increased statistical power.

To conclude, I have identified a protective variant rs1990621 in TNEM106B associated with increased neuronal proportion through bulk RNA-Seq deconvolution and cell type proportion QTL analysis. This observation also replicated previous findings of the protective variant rs1990622 in FTD risk, which is in high LD with rs19990621[266]. Besides, I also observed the C allele of rs429358 (codetermine APOE ε4 isoform with rs7412 C allele) associated with decreased neuronal proportion as it was hypothesized. It suggested potential involvements of both APOE and TNEM106B in neuronal protection mechanisms underlying neurodegenerative and normal aging processes, and supported previous observation of interactions between these two genes[146] in AD cohort. I speculate that TNEM106B related lysosomal changes might be involved in the common pathway underlying neuronal death and astrocytosis in neurodegenerative disorders and normal aging cohorts. With larger sample size and higher deconvolution resolution, this approach will reveal more biologically relevant and novel loci associated with changes in cellular composition that are important for understanding both disease etiology and healthy aging.
Chapter 4: System biology approaches revealed transcriptomic profiles of *TREM2* and *PSEN1*
4.1 Abstract

**Background:** Using network analysis approaches, previous studies had revealed several hub gene or pathways that were verified or later identified as key players in disease etiology underlying AD. In sporadic AD, previous studies from the lab have identified *MS4A* gene cluster significantly associated with soluble TREM2 level in CSF. However, from GWAS result it was unclear which *MS4A* gene is the key regulator of *TREM2*. In autosomal dominant AD, mutations in the *APP, PSEN1* and *PSEN2* genes and lead to familial early onset AD. However, the downstream pathogenic events triggered by these risk and pathogenic variants are still not fully understood. By employing an integrative network approach, I aim to more accurately identify which gene is the key regulator of *TREM2* in sporadic AD and the downstream genes and pathways altered by *PSEN1* mutation in autosomal dominant AD.

**Methods:** To determine which one of the *MS4A* genes are implicated in TREM2 biology, I employed alternative approaches to explore gene regulatory networks from RNA-Seq data. To identify causal genes under the genomic locus identified by the CSF TREM2 GWAS, I combined weighted correlation network analysis (WGCNA) method to identify a module that includes *TREM2* co-expressed genes. Then I used Bayesian network inference to learn causality. To study the downstream effects of Mendelian mutations in *PSEN1* associated with Autosomal Dominant Alzheimer’s disease, I applied a seed-based approach to study the genes that are significantly co-expressed with *PSEN1*, and constructed gene networks using WGCNA. This analysis includes both *PSEN1* mutation carriers, non-carriers and neuropathological-free controls. The network was annotated with gene differential expression, cell type information, and functional pathway analysis.
**Results:** My analysis indicated that *MS4A4A* and *MS4A6A* are in TREM2 module, and inferred that *MS4A4A* is the key regulator of *TREM2*. For the downstream events of the Mendelian gene *PSEN1*, I identified 47 genes only present in control cohort that were potentially disrupted in *PSEN1* mutation carriers; I also found 13 genes only present in *PSEN1* mutation carriers but not in control cohort that are potentially acquired as downstream transcriptional events altered by *PSEN1* mutations. Among them, I highlighted the genes *LMNA*, *DOCK1*, and *DYNC1LI2* and discussed them in detail, that were previously associated with Alzheimer’s Disease.

**Conclusions:** My study demonstrated the potential of using both system-based and seed-based network approaches in replicating and discovering AD related genes and their interactions. In sporadic AD cohort, I identified *MS4A4A* might be a key regulator for *TREM2*. In autosomal dominant AD cohort, I identified total of 60 genes that are lost or acquired in the *PSEN1* associated pathways.
4.2 Introduction

4.2.1 From polygenic to omnigenic - a network interpretation

To tackle disease etiology, one important theme for diseases with genetic components is to figure out how genetic variants explain phenotypic variability. A monogenic theory inspired by Gregor Mendel’s work states that one disease could be explained by one mutated gene following a Mendelian inheritance pattern. Examples are familial early onset neurodegenerative disorders such as autosomal dominant AD, which can be explained by rare mutations in one single gene that results in high disease penetrance. However, diseases with complex traits, for example late onset sporadic AD, do not follow this pattern. GWAS performed in sporadic AD studies have identified dozens of variants across the genome, and many of those are common variants with low to medium effects (Figure 1.2)[148]. This pattern is more similar to quantitative genetics inspired by Ronald Aylmer Fisher’s infinitesimal model that a quantitative trait is influenced by an infinitely large number of genes. Accumulations of large number of common variants within multiple genes explain much of the heritability. These polygenic effects together with complex interactions with the environment are often observed in diseases with complex traits. Based on empirical evidences, the polygenic model has later been expanded to an “omnigenic” model for complex traits[31]. They proposed that core genes may have strong and direct effects on disease risk, but they only account for a small portion of total heritability. Any variants that have disease relevant tissue specific effects may contribute nontrivial effects on disease risk. The variants may exert their effects on core genes through highly interconnected gene regulatory network such that the collection of the small effects together explain the missing heritability[31].
4.2.2 Network analysis as a powerful tool

Network biology[21, 22, 269] has demonstrated great success in understanding biological systems and identifying disease related factors. With large scale data collection and advancement in computational powers, various types of networks have been proposed to capture the interactions among different elements (zoom-in view) and to gain a systematic view of the topological and dynamical properties of a biological system (zoom-out view). On a gene level, gene regulations can be depicted as gene regulatory networks as mentioned above in the omnigenic model. The modular and hierarchical organization of gene regulatory networks captures the information flow from regulators to their binding sites. On an RNA level, microarray and RNA-Seq technologies have enabled generations of transcriptome-wide gene co-expression networks to understand associations among transcripts and gene expression synchronicity. On a protein level, yeast two-hybrid screens are able to identify protein-protein interactions in vivo[89]. On a cellular level, brain neural networks generated from tracer injections have captured the topology of neural signaling highways underlying cognitive functions[85]. On a system level, networks generated from functional connectivity MRI facilitate understanding of how different parts of the brain segregated into functional modules and communicating intrinsically without explicit task being performed[213]. On a disease level, human disease interactome has been proposed to identify shared pathways among known or unknown comorbidities that shed lights on drug repurposing[50, 183]. On a population level, social networks could help identify key features of infectious diseases, such as risks for acquisition and effective interventions, through learning social aspects of disease transmission.
While offering the capability of modeling any type of biological interactions at different levels, network analyses provide a unique and powerful approach that can combine elements or layers from different modalities and produce integrated models to study interactions among multiple networks. One example of an integrative network analysis of combining obesity and social network showed that the social ties among friends have a larger effect on obesity risk than genetic risk factors (Figure 4.1)[19, 52].

![Network medicine from obesity to the "diseasome". Reproduced with permission from Barabasi[19]. Copyright Massachusetts Medical Society.](image)
4.2.3 Basic concepts in gene co-expression networks

In biological systems, networks are defined as a collection of biological elements and interactions among them. In the context of a gene co-expression network, nodes are genes and edges are relations between the expression of a pair of genes, such as Pearson’s correlation. If the network is unweighted, an edge means the correlation between a gene pair is above a hard cut-off, and there will be no edge if the correlation is below the cutoff. If the network is weighted, the edge represents weighted magnitude of correlation between a gene pair. In an undirected network, such as results from WGCNA, there is no directionality in the connection between two nodes. In a directed network, such as directed acyclic network generated from Bayesian network inference or structural equation modeling (SEM), information flow will be depicted as directed arrow pointing from parent node to child node. This chapter focuses on static and deterministic networks that capture network structural topology from a single RNA-Seq snapshot when RNAs are extracted from the biological system. Dynamic and stochastic networks can be generated with temporal data and data with biological noises. Causal network can also be generated with intervention data.

Regarding the global topology of a network, three common types of networks are showed in Figure 4.2. Two important metrics to describe network property are degree distribution and adjacency matrix. Degree is the number of edges a node has with other nodes (denoted as k), and degree distribution is the probability distribution of these degrees over the whole network denoted as p(k). Adjacency matrix is used to represent whether a pair of nodes are connected. It is a matrix with 0 and 1; value 1 represents the nodes are connected and adjacent to each other, and 0 represents that the nodes are unconnected. Undirected graph adjacency matrix is symmetrical and directed graph adjacency matrix is unsymmetrical. In a random network, the
degree distribution $p(k)$ follow a Poisson distribution, and connected nodes are randomly distributed in the adjacency matrix. Network path is another important metric, which measures how many steps required to connect two nodes in the network, and minimal number of steps is called the shortest path length. One important property of a random network is that the shortest path length is much smaller than a regular network. Many theoretic and functional works in network science are based on random network, however, it has two limitations – first, it does not have local clustering structure with its randomly distributed connections depicted in adjacency matrix; second, the degree distribution of nodes following a Poisson distribution do not account for the formation of hubs that mostly observed in real world networks. To resolve the first limitation of lacking local clustering, a Watts-Strogatz model has been proposed to generate graphs with small-world property by rewiring edges from a regular lattice network with random probabilities[277]. This random rewiring process created long-range connections with small path length like a random graph while retaining high local clustering properties of a regular network. Thus, the degree distribution of small world network is similar to a random network but with high local clustering property shown in adjacency matrix. Later a scale free model is proposed based on empirical evidence of real world networks including World Wide Web and citation patterns in science[20]. This model explained the hubs observed in many real-world networks that a random network model does not explain. They found that large scale complex networks exhibit a high degree of self-organizing phenomena that networks expand by adding new vertices to already well established highly connected vertices, which explained how hub nodes emerged from chaos. In a scale-free network, the degree distribution of nodes decays as a power law. This feature holds true in any scales of the network – hence the name “scale-free” network. Rigorous
modeling is required to examine the power law distribution of empirical data to determine if a network fulfills the criteria[53].

Figure 4.2 Random, small-world, and scale-free network properties. A) random network with 73 connections among 20 nodes assigned randomly; B) Small-world network with high local clustering and short average path lengths with ‘hub and spoke’ architecture; C) scale-free network with ‘hub and spoke’ architecture maintained at multiple spatial scales. Image reproduced from Stobb et al.[251] with permission.

Figure 4.3 Small-world network generated from Watts-Strogatz model. Image from Watts et al.[277] with permission.
4.2.4 Network analysis in Alzheimer’s disease

Gene co-expression network analysis in late-onset AD cohorts had revealed several hub gene or pathways that were verified or later identified as key players in disease etiology[184, 287]. For example, a system based approach using WGCNA had been applied to microarray data derived from prefrontal cortex tissues collected from LOAD patients and 173 non-demented healthy controls. They identified an immune response related module, which contains an important AD risk gene TREM2. Rare variants in TREM2 have been found to be associated with sporadic AD risk with moderate effect[114, 144, 148]. The rare TREM2 variant p.R47H (rs75932628) carriers exhibit increased AD risk by a range from 1.7-fold to 3.4-fold[112, 212]. In the TREM2 module identified from network analysis, Zhang et al. focused on TYROBP(DAP12) gene which was identified as an adapter protein for TREM2[30] (Figure 4.4).

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Figure 4.4 TREM2 module identified in LOAD cohorts. A module enriched for immune function and pathways contains TREM2, TYROBP, and MS4A gene clusters. Image from Zhang et al.[287] with permission.
Our lab has recently identified that the *MS4A* gene cluster is a key regulator of soluble TREM2 in CSF[70]. Using GWAS of a large number of subjects (N = 813) we identified a locus in chromosome 7 that shows significant association with CSF TREM2 levels (rs1582763; p=1.15 \times 10^{-15}), and replicated in an independent dataset (N = 580). Several members of *MS4A* gene cluster, *MS4A4A* and *MS4A6A* are tagged by this signal, which makes difficult the identification of the causal *MS4A* gene affecting CSF TREM2 levels. I envision that transcriptomic data would provide additional orthogonal evidence. To identify the relationship between TREM2 and MS4A gene cluster, I performed a system-based approach on sporadic AD cohort and focused on the module containing *TREM2* in the first part of this chapter.

In the second part of this chapter, I focused on the study of the transcriptomics downstream analyses of genetic mutations causal of autosome dominant AD (ADAD). Mutations in the *amyloid-beta precursor protein* (*APP*) and presenilin genes (*PSEN1* and *PSEN2*)[43] cause ADAD which is typically associated with Mendelian inheritance pattern and early-onset (30 ~ 50 years old) disease symptoms. Although the disease phenotype may be a consequence of an abnormality in a single effector gene product particularly in the Mendelian form of AD, given the highly-interactive functional crosstalk within biological organism and the complex disease etiology of AD, this dysregulation by a single gene may intertwine with various pathological processes and altered downstream events that interact in a complex network. To investigate the etiological heterogeneity of AD, I performed a seed-based network analysis in an ADAD cohort, carriers of *PSEN1* gene mutation, to identify altered downstream transcriptional events triggered by this ADAD gene. By annotating the network with differential gene expression, cell type information, and functional pathway analysis, I identified some target genes and related
pathways that were either disrupted or emerged in PSEN1 mutation carriers through network analysis.

4.3 Methods

4.3.1 Samples

**TREM2 Study**

I accessed (AMP-AD portal synapse ID = 3157743) RNA-Seq data from 219 AD and non-demented control brains from Mount Sinai School of Medicine (MSSM) ascertained from four cortical regions: anterior prefrontal cortex (APC), inferior frontal gyrus (IFG), superior temporal gyrus (STG), and parahippocampal gyrus (PHG) (Table 4.1). Data retrieval and collection of MSSM, Knight-ADRC, and DIAN have been documented in detail in Chapter 2.

<table>
<thead>
<tr>
<th>Table 4.1 TREM2-MS4A Study Demographic</th>
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<tr>
<td>N</td>
</tr>
<tr>
<td>----</td>
</tr>
<tr>
<td>MSSM</td>
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</tbody>
</table>

**PSEN1 Study**

RNA-Seq was generated for 15 *PSEN1* carrier brains from The Dominantly Inherited Alzheimer Network (DIAN) and 14 non-demented controls from The Charles F. and Joanne Knight Alzheimer's Disease Research Center (Knight ADRC)[153]. We identified three additional participants from the Knight-ADRC study with *PSEN1* (p.A79V, p.I143T, p.S170F) mutations (Table 4.2).

I accessed (AMP-AD portal synapse ID = 3157743) RNA-Seq data from 67 non-demented control brains from Mount Sinai School of Medicine (MSSM) ascertained from four
cortical regions: anterior prefrontal cortex (APC), inferior frontal gyrus (IFG), superior temporal gyrus (STG), and parahippocampal gyrus (PHG).

GEO replication data was accessed from GSE39420, which is collected from 14 patients (7 sporadic EOADs and 7 monogenic familial ADs with PSEN1 mutation) and 7 neurologically healthy controls. Samples were hybridized in a Human Gene 1.1 microarray from Affymetrix[13].

<p>| Table 4.2 PSEN1 Study Demographic |</p>
<table>
<thead>
<tr>
<th>ZEN</th>
<th>N</th>
<th>Age</th>
<th>% Male</th>
<th>% ApoE4+</th>
<th>Control</th>
<th>PSEN1</th>
<th>EOAD</th>
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<tr>
<td>MSSM</td>
<td>67</td>
<td>80.1 ± 8.39</td>
<td>44.8</td>
<td>10.4</td>
<td>67</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Knight-ADRC</td>
<td>17</td>
<td>82.3 ± 18.5</td>
<td>35.3</td>
<td>17.6</td>
<td>14</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>DIAN</td>
<td>15</td>
<td>49.1 ± 7.14</td>
<td>66.7</td>
<td>14.3</td>
<td>0</td>
<td>15</td>
<td>0</td>
</tr>
<tr>
<td>GEO</td>
<td>21</td>
<td>55.6 ± 7.65</td>
<td>76.2</td>
<td>14.3</td>
<td>7</td>
<td>7</td>
<td>7</td>
</tr>
</tbody>
</table>

4.3.2 Data processing and quality control

TREM2 Study

Data QC and preprocessing of MSSM dataset have been documented in Chapter 2.3.2. MSSM cases and controls gene expressions were derived from Star alignment to human GRCh37 primary assembly and quantification using --quantMode TranscriptomeSAM. Because low expressed genes tend to reflect noise and produce insignificant correlation, I removed genes with gene counts less than 4 in more than 75% subjects. To normalize gene expression respect to library size, regularized logarithm transformation was applied to raw counts of the gene expression using rlog function from DESeq2 R package. ComBat function was applied to the data to remove potential batch effect. After which, a linear regression model was applied to regress out covariate on a per-gene basis. Covariates factors included in the model are PC1, PC2,
PC3 inferred from genomic PCA analysis to account for ethnic stratification; sex, age at death, post-mortem index, RIN to account for general demographics and RNA-Seq tissue pre-sequencing quality; and RNA-Seq post-sequencing metrics such as ribosomal contents, mapped reads number (uniquely mapped and multi-mappers that mapped to multiple loci) to account for alignment performances. The residuals from the linear regression were used as inputs to infer gene expression correlation. Apart from gene expression values, cognitive performance measurement CDR and Tau pathology load measurement Braak staging values were also added for computing correlation, from which I could infer what genes are closely correlated with these clinical and pathological traits. Besides, cellular composition inferred from the dataset as documented in Chapter 2 deconvolution procedure were also added to later correlation computing and network construction. Cell type information inferred from the data using deconvolution method were also added as nodes. For each region, the top third most variable genes were selected. A joint set of the top genes from all four regions accounted for a fifth of the whole transcriptome genes. The top fifth most variable genes together with CDR, Braak staging, and cell type proportion were selected to run multi-tissue correlation for later network construction.

PSEN1 Study

Data QC and preprocessing of MSSM, Knight-ADRC, and DIAN have been documented in Chapter 2.3.2. For the RNA-Seq data from MSSM gene expression were performed similarly as the TREM2-MS4A study as described above, the only difference is that only non-demented controls from MSSM were used to construct the network to avoid the confounding factors from dramatic neuronal loss or astrocytosis in ADAD as I observed and documented in Chapter 2.
For microarray data from GSE39420, probes were mapped to corresponding genes, and each gene expression value was derived from averaging probe expression values for each gene. Each gene expression level was adjusted for subject sex, age at death, and post-mortem interval hours by fitting a linear regression model and the residuals were derived for use in downstream analyses. These covariate adjustment procedures were performed for control, FAD-PSEN1, and EOAD separately.

4.3.3 Repeated measures correlation

I accessed RNA-Seq data MSSM ascertained from four cortical regions: APC, IFG, STG, and PHG. Because there is more than one tissue collected from the same subject, the assumption of independent observations when applying standard correlation methods is violated. To aggregate data collected from multiple tissues in MSSM, I integrated the measures from these four brain regions by running repeated measures correlation tool (rmcorr R package)[18] to calculate repeated measures correlation (rmcorr) of the MSSM controls, which is a statistical technique to determine the overall within-individual relationship among paired measures assessed on two or more occasions.

4.3.4 Network construction

*TREM2* study system-wide network construction

With a system-wide approach, Weighted Gene Co-Expression Network Analysis (WGCNA)[164] was applied to the transcriptome rmcorr matrix derived from MSSM AD cases and control subjects. WGCNA enforces the connectivity to exhibit a power-law distribution. This power-law distribution renders a scale-free topology to the network, which will be applied to the top fifth most variable genes across all four regions. Both dynamic tree cutting and static tree
cutting with signed networks and minimum module size 200 were used, but it has been shown that the dynamic tree cutting and signed network might be more biologically relevant compared to static tree cut and unsigned network[289]. Raising the correlation to a power will help reduce the noise of the correlations in the adjacency matrix. To select the appropriate power value, pickSoftThreshold function[288] from the WGCNA package was applied to the rmcorr matrix to select the power when the network most resemble a scale-free graph while keeping the highest network connectivity.

*TREM2* study Bayesian network construction

To infer probabilistic relationships between the nodes in a network module, Bayesian network analysis was applied to the module derived from the system-wide WGCNA results that contains *TREM2* gene. Normalized and covariate adjusted gene expression matrix from parahippocampal region of MSSM dataset was used, and only genes in the same module with *TREM2* were included in the Bayesian network construction. Using the discretize function from bnlearn R package[193], the continuous gene expression data were transformed into quantiles for later network structure and parameter learning. To infer or measure the degree of confidence of arc strength of Bayesian network, 200 nonparametric bootstrap iterations were applied to the data to estimate the relative frequency (strength) of every possible arc[94] using the boot.strength function implemented in bnlearn package. Arcs with strength more than 0.38 and probability of direction more than 0.5 were kept, and final network was derived from averaging across 200 bootstraps. The network was plotted using graphviz.plot function with highlighted v-structure arcs. Genes related to AD risk and genes related to top pathway results (adjusted p-value less than 0.01) were color coded respectively.
**PSEN1 study seed-based network construction**

Using a seed-based approach, I pre-selected genes that are co-expressed with PSEN1 to build a co-expression network. The genes that are correlated with PSEN1 expression are potentially linked to AD pathology by assuming ‘guilt-by-association’. In this seed-based approach, I used PSEN1 gene as a bait to expand to genes that co-expressed/correlated with PSEN1 in PSEN1 variant carriers and/or healthy controls.

To build upon the MSSM controls co-expression network, I first selected genes that are significantly correlated with PSEN1 in the MSSM control rmcorr results at correlation p-value < 0.05. Then the genes from parietal PSEN1 carriers and healthy control subjects that overlapped with the significant correlated gene in MSSM controls were selected following the two criteria: (1) The gene correlation with gene PSEN1 in parietal dataset has to fall into the 95% confident intervals of rmcorr results of the MSSM controls; (2) The correlation direction in parietal dataset has to be congruent with the direction of the MSSM controls rmcorr correlation.

**4.3.5 Network robustness evaluation**

By applying a bootstrapped version of WGCNA (rWGCNA)[100], it will reduce potential bias introduced by outlier samples. I performed 50 iterations of network construction with randomly selected 66.66% of the total samples. The resulting 50 networks will be merged into one large, final consensus network. The robustness of the networks was evaluated by comparing those to the final consensus network.
4.3.6 Network functional annotation

Differential gene expression analysis using DESeq2 R package, and various pathway analysis and gene enrichment analysis were applied to annotation the genes in the network, such as Kyoto Encyclopedia of Genes and Genomes (KEGG), Gene Ontology Enrichment Analysis (GO term), and PANTHER implemented in Enrichr online interactive tool[47, 158]. Cell type composition derived from deconvolution, CDR, and Braak staging were labeled in the network to examine cell-type specific enriched modules and modules related to cognitive and pathological measurements. To annotate the genes, the list of interested genes were queried with FUMA’s GENE2FUNC online tools[276] to investigate tissue specificity, reported GWAS catalog genes, TF targets, microRNA targets, and etc.

Differential gene expression and correlation analyses were repeated in the independent PSEN1 dataset (GSE39420) collected from 14 patients (7 EOAD and 7 FAD-PSEN1) and 7 neurologically healthy controls.

4.4 Results
4.4.1 Study design

TREM2 Study

For the TREM2 study, both AD sporadic cases (N = 170) and cognitive normal controls (N = 49) were included to build the network (Table 4.1). Tissues from four brain regions (Figure 4.5) were collected and followed by RNA-Seq data generation. The same extensive QC and quantification processes were applied as documented in Chapter 2. Repeated measure correlation was applied to gene expression quantification results from four regions to derive a gene expression correlation matrix. With a system-based approach, top 1/5 most variable genes
across the four regions were used to construct networks with scale free topology using WGCNA. The module containing TREM2 gene was further analyzed with Bayesian network inference and functionally annotated with pathway analysis.

**PSEN1 Study**

As discussed in **Chapter 2**, I observed brain carriers of pathogenic mutations in APP, PSEN1 or PSEN2 presented lower neurons and higher astrocytes relative proportions compared to sporadic AD and controls. With such extensive neurodegeneration, I concerned that the network built with PSEN1 mutation carriers would be confounded by the destructive consequence of neurodegeneration. To investigate early transcriptional events trigged by pathogenic mutations in PSEN1, I built the network with cognitive normal controls and genes correlated with PSEN1 using a seed-based approach. First, I selected genes that are correlated with PSEN1 in both cognitive normal controls and PSEN1 mutation carriers. Then I built a network based on correlations of these selected genes in controls across four different brain regions in MSSM to capture a control network topology. Then nodes were annotated in regard to its correlation with PSEN1 in either controls or PSEN1 mutation carriers to infer their functional roles in early disease progression. I hypothesized that network modules enriched with genes correlated with PSEN1 in mutation carriers could be involved in early transcriptome changes in PSEN1 mutation carriers.
Figure 4.5 Study design. For TREM2 study, network was constructed from MSSM AD sporadic cases and controls. For PSEN1 study, network topology was built from MSSM control participants and functional modules were annotated in related to PSEN1 variant carriers and healthy controls.
4.4.2 *MS4A* gene clusters are associated with *TREM2* in AD sporadic and control network analysis

**TREM2 Study WGCNA network construction**

The 20% of most variable genes from four regions of AD sporadic cases and controls were selected to construct the network using WGCNA. The network connection or topology was built upon gene co-expression patterns. Genes that are connected in a co-expression network are correlated or synchronized in their expression pattern[97, 223]. In a gene expression correlation matrix, any pair of genes would yield a non-zero value, which means any gene pair could be correlated to some extent and perfect independency does not exist. Thus, the challenge is how to determine the threshold at which two genes are considered as co-expressed. An intuitive way to solve this problem is to select a value as a hard cut-off. A binary value is assigned to each gene pair; it will be 1 if the gene pair correlation is above the threshold or 0 if it is below the threshold. This binary scenario may occur in certain biological systems such as neuron firing, but it does not fit gene expression patterns. Rather than this hard thresholding approach, the WGCNA researchers proposed a soft thresholding approach that assigns a weight to each gene pair to derive a weighted gene co-expression network. The network also bears a scale-free topology, which has been shown to be more biologically relevant by both theoretical and empirical evidences[288]. The property of scale-free network is defined as the probability of a node that is connected with k other node decays as a power law. This type of network contains a few hubs with high degree of connectivity compared to the vast majority of non-hub nodes with low connectivity. The network is highly resistant to attacks on non-hub nodes, but removing hub nodes will be deleterious or change the network topology dramatically. The hub nodes are essential for survival in biological systems proved by yeast protein network studies[118, 142].
To build a scale free network using WGCNA, taking my study as an example, the first step was to define gene co-expression similarity using repeated measure correlation derived from gene expression of four brain regions. Compared to all the other network analysis that use single region transcriptional data to build a region-specific network, this study integrated data from four brain regions to build a network representing the cerebral cortex. Repeated measure correlation was applied instead of Pearson correlation because the four cortical regions are not independently collected, thus the assumption of independent observations when applying standard correlation methods was violated. Then the similarity (correlation) matrix was transformed into an adjacency matrix with appropriate power parameters fitting the power adjacency function[288]:

\[ a_{ij} = \text{power}(s_{ij}, \beta) \equiv |s_{ij}|^{\beta} \quad \text{(Equation 4.1)} \]

As discussed above, the scale-free network nodes follow the power law:

\[ p(k) \sim k^{-\gamma} \quad \text{(Equation 4.2)} \]

To define a scale-free topology criterion, \( R^2 \) is the model fitting index of the linear model that regresses \( \log(p(k)) \) on \( \log(k) \) based on Equation 4.2, and higher \( R^2 \) means a better model fit to the scale free topology. \( R^2 \) increased as power increased (Figure 4.6A), thus higher power would yield a network more resembling scale free topology. However, there was also a tradeoff between scale free network resemblance and connectivity (Figure 4.6B), and too sparse networks forfeit too much connection information. In order to construct a network with scale-free topology and reasonable connectivity, the first power value that passed \( R^2 = 0.8 \) was picked to fulfill scale-free topology criterion while retaining high enough connections to investigate
nodes relationships. To detect modules in the co-expression network, the topological overlap dissimilarity was measured and a topological overlap matrix (TOM) was derived to reflect relative interconnectedness between two nodes. Based on TOM, hierarchical clustering and different tree-cutting approaches were applied to determine module boundaries and gene memberships[165]. Figure 4.7A showed different modules derived from dynamic tree cutting on hierarchical clustering of the transcriptome wide TOM matrix containing the top 20% most variable genes. The dynamic tree cutting is a top-down approach that interactively decomposes and combines cluster branches until the assignment of module becomes stable. One problem of this dynamic tree cutting method is that it may fail to assign some tree branches, although it was not an issue in my TREM2 study (Figure 4.7A). To resolve this potential problem, Figure 4.7B showed another module assignment of the same hierarchical clustering tree with a dynamic hybrid cutting method, which is a bottom-up approach that improves the detection of any unassigned branches. Noticeably, for the module I was interested in containing TREM2 gene (Figure 4.7AB labeled with star) the two tree cutting methods produced almost identical assignments with only 12 (out of 456) more genes in the hybrid tree cutting. The TREM2 modules derived from both methods showed almost identical gene memberships and network topology (Figure 4.7CD). TREM2 gene is expressed in microglia and involved in immune responses in AD. Not surprisingly AD risk related genes HLA-DRB1 and HLA-DRB5, which play central roles in immune system by presenting peptides derived from antigens, were also present in the same module with TREM2. Another sporadic AD risk gene in this module is DSG2, which is important for cell to cell adhesion functions, and its cytoplasmic domain anchors the cytoskeleton by interacting with plaque proteins in the desmosome-intermediate filament complex[60]. What really caught my attention is the microglial MS4A gene cluster that was
located in the same co-expression module with *TREM2*, because we have recently observed that common variants in the *MS4A* region were significantly associated with elevated CSF soluble TREM2 level (rs1582763; p-value = 1.15×10^{-15})[70].

**Figure 4.6** TREM2 network soft thresholding. A) $R^2$ is the scale-free model fitting index. Higher $R^2$ means a better model fit to the scale free topology. $R^2$ was plotted for power values ranging from 1 to 20. A $R^2 = 0.8$ cut-off line was plotted. B) However, there was also a tradeoff between scale free network resemblance and connectivity as the mean connectivity decreases as the power increases.
In the previous study of network analysis in LOAD using WGCNA, Zhang et al. identified an immune/microglia module from prefrontal cortex microarray data. In this module containing \( \text{TREM2} \), \( \text{TYROBP} \) scored the highest based on both regulatory strength and differential expression, which is an adaptor protein of TREM2. Apart from these two genes, \( \text{MS4A4A} \) and \( \text{MS4A6A} \) were also located in the same module. Although I identified \( \text{MS4A4A} \) and \( \text{MS4A6A} \) in the same module in my analysis, I seemed to have missed an important finding from Figure 4.7 TREM2 network TOM clustering and gene module assignment. A) Dynamic tree cutting B) Dynamic hybrid cutting. C) TREM2 module derived from dynamic tree cutting. D) TREM2 module derived from dynamic hybrid cutting. The module contained TREM2 is starred.

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Zhang et al., which is *TYROBP*. Apart from the fact that my data is generated from RNA-Seq that is different from their microarray data, my analysis also aggregated samples from non-independent multiple tissues through repeated measures correlation. To recapitulate their finding and to understand why I missed *TYROBP* in my *TREM2* module, I performed a robust WGCNA analysis on single tissue region from anterior prefrontal cortex (BM10), which is the same region where they identified the *TREM2-TYROBP* module. The robust WGCNA is a resampling version of a regular WGCNA that I randomly resampled two thirds of the total sample size for each iteration, and repeated the process for 50 iterations to assess module assignment robustness. In this analysis, I focused on four genes that are *TREM2, MS4A4A, MS4A6A*, and *TYROBP*. *MS4A4A* and *MS4A6A* are from the same *MS4A* gene cluster located nearby on chromosome 11. Their expression patterns are highly correlated so they are included in this robustness assessment as positive controls. In the full run with all available samples from BM10 (N = 181), *MS4A4A, MS4A6A*, and *TYROBP* were assigned into module 12, colored as tan module in Figure 4.8 labeled with star. *TREM2* was not assigned to any module in the full-size run. It worth mentioning that in the previous study, Zhang et al. also applied Bayesian network analysis and pathway analysis to further annotate the submodules from their *TREM2-TYROBP* network. In their pathway analysis, *MS4A4A, MS4A6A*, and *TYROBP* were segregated into the complement pathway, whereas *TREM2* was assigned to a separate pathway named Fc receptor system. This observation suggested that compared to *TREM2-TYROBP* co-expression, there might be a stronger relationship within *MS4A4A-MS4A6A-TYROBP* co-expression pattern. My robust WGCNA results supported this hypothesis but more in-depth simulations with 1000 runs are required to further validate this hypothesis.
Figure 4.8 Robust WGCNA of anterior prefrontal cortex. Top hierarchical tree is derived from full dataset with all the subjects. Two thirds of the total sample size were randomly sampled for each resampling run. One full size run and total of 50 resampling runs with consensus module assignment were shown as color bars. The module contained $TREM2$ is starred.
### Table 4.3 Robust WGCNA simulation of TREM2 module

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TREM2 Study Bayesian network construction

To further infer the relationship between *TREM2* and *MS4A*, Bayesian network inference was applied to the WGCNA module containing *TREM2* gene. Bayesian network infers factorized probability distribution from gene expression, and incorporates it into directed acyclic graphical representation of the network. As shown in Figure 4.9, compared to undirected graphs from WGCNA, Bayesian network inference produces directed graphs with arrows pointing from parent nodes to child nodes. Directed acyclic graph has no directed cycles, which generates a topological ordering of nodes. The resultant network was produced from averaging across 200 bootstraps with both connection strength and direction probability hard cutoffs. In the averaged final network, there are numerous singletons, doubletons, and small branches with less than 10 nodes. The large central branch containing *TREM2* was labeled with blue, and previously identified AD risk genes were labeled with red. It worth noticing that there was a gene regulatory cascade from *MS4A6A* to *MS4A4A* to *TREM2*, suggesting a potential regulatory effect of *MS4A* genes on *TREM2*. This observation has been supported by our recent *in-vitro* MS4A knockdown study using cultured macrophage[70]. We observed that the soluble TREM2 level in cell culture was decreased after *MS4A4A* knockdown but it does not respond to *MS4A6A* knockdown. The knockdown and Bayesian network results together suggest *MS4A4A* might be the direct regulator of *TREM2*.

Pathway analysis was performed to further annotate the function of this TREM2 module, and top 8 functional pathways were labeled in the graph. The central large branch containing *TREM2*, *MS4A* genes, and *HLA-DRB* genes were involved in immune system related pathways, such as bacterial or parasitic infections, autoimmune disease systemic lupus erythematosus, toll-
like receptors mediated pathogen recognition, phagocytosis and complement cascades. The second largest branch next to the blue branch was enriched with ribosomal pathway. Compared to the pathway analysis results from Zhang et al. TREM2 module, my top 8 functional pathways with many overlapped genes replicated 4 out of 5 pathways that they highlighted for their TREM2 network.

**TREM2–MS4A Gene Network**

AD related genes
- Ribosome
- Staphylococcus aureus infection
- Complement and coagulation cascades

Phagosome
- Systemic lupus erythematosus
- Tryptophan metabolism
- Toll-like receptor signaling
- Malaria

Figure 4.9 TREM2-MS4A Gene Network. AD related and top eight pathways are color coded accordingly. The largest branch is color coded in blue. V-structures are labeled in bold.
4.4.3 Disrupted and acquired genes identified in network module of \textit{PSEN1} mutation carriers

\textbf{PSEN1 study WGCNA network construction}

Similar to the TREM2 study, I accessed RNA-Seq data from four cortical regions of MSSM[3], and estimated the gene expression correlation combining all four areas using repeated measure correlation. There are two key methodological differences in the approach I employed to model PSEN1 network: first, only non-demented controls were included to construct the network; second, instead of a system-based approach selecting most variable genes from the whole transcriptome, only genes correlated with \textit{PSEN1} expression pattern were included. The genes that are correlated with \textit{PSEN1} expression are potentially linked to AD pathology by assuming ‘guilt-by-association’. With this seed-based approach using \textit{PSEN1} as a bait, I pre-selected genes that are co-expressed with \textit{PSEN1} gene in both \textit{PSEN1} mutation carriers (N = 18) and non-carrier non-demented controls (N = 14) to build a co-expression network (Figure 4.10). Collectively, total of 5,809 genes correlated with \textit{PSEN1} were selected from \textit{PSEN1} non-carriers (Figure 4.10A) and mutation carriers (Figure 4.10B) that were jointly overlapped with genes correlated with \textit{PSEN1} from MSSM non-demented control participants.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure.png}
\caption{PSEN1 seeded network gene selection. Select genes from \textit{PSEN1} mutation carriers and non-carriers correlated with MSSM control \textit{PSEN1} gene expression.}
\end{figure}
With similar soft thresholding power selection and hierarchical clustering tree cutting approaches, modules containing \textit{PSEN1} were produced for both dynamic tree cutting (Figure 4.11AC) and dynamic hybrid cutting (Figure 4.11BD). Compared to the dynamic tree cutting module (Figure 4.11A), the dynamic hybrid cutting (Figure 4.11B) lost branches to the left of the module and gained branches to the right of the module labeled by arrow heads. Based on a human visual inspection of the hierarchical clustering tree, the gain of the right branches might be reasonable but the loss of the left branches seems to be spurious. However, the dynamic hybrid cutting subdivided the \textit{PSEN1} module and grouped the leftmost branches to the large module on the left of \textit{PSEN1} module (Figure 4.11B turquoise module), due to which the dynamic hybrid cutting \textit{PSEN1} module lost \textit{BACE1}, \textit{BIN1}, and oligodendrocyte proportion measures (Figure 4.11D) compared to dynamic tree cutting (Figure 4.11C). Based on queries of \textit{PSEN1} in Brain RNA-Seq database of human brain purified cell type specific expression analysis\cite{291} (http://www.brainrnaseq.org/; Figure 4.12A) and our single nuclei RNA-Seq dataset\cite{66} (http://ngi.pub/snuclRNA-seq/; Figure 4.12B), \textit{PSEN1} is mostly expressed in oligodendrocyte, which support dynamic tree cutting \textit{PSEN1} module assignment.
Figure 4.11 PSEN1 network TOM clustering and gene module assignment. A) Dynamic tree cutting B) Dynamic hybrid cutting. C) PSEN1 module derived from dynamic tree cutting. D) PSEN1 module derived from dynamic hybrid cutting. The module contained PSEN1 is starred.

Figure 4.12 PSEN1 is mostly expressed in oligodendrocyte. A) PSEN1 expression in purified cell type specific expression analysis[291]. B) PSEN1 expression in our single nuclei RNA-Seq dataset[66].
With the dynamic tree cutting $\textit{PSEN1}$ module, I performed differential correlation and differential expression analysis and functional annotation to identify genes that are altered in $\textit{PSEN1}$ mutation carriers. The analysis was replicated in another independent $\textit{PSEN1}$ mutation dataset. The genes were categorized into five groups and depicted as a Venn diagram[120] to show the overlaps among the groups (Figure 4.13). In group A, there were 268 genes that are significantly correlated with $\textit{PSEN1}$ expression in Knight ADRC non-demented controls. In group B, there were 264 genes that are significantly correlated with $\textit{PSEN1}$ expression in Knight ADRC $\textit{PSEN1}$ mutation carriers. In group C, there were 269 genes differentially expressed significantly comparing Knight ADRC $\textit{PSEN1}$ mutation carriers to non-demented controls. In group D, there were 25 genes that were differentially correlated with $\textit{PSEN1}$ when comparing correlation derived from group A to group B[73]. In group E, there were 64 genes specifically replicated in GEO $\textit{PSEN1}$ mutation carrier data (removing genes replicated in GEO $\textit{PSEN1}$ non-carrier early onset AD cohort). Genes from combinations of A + C, A + D, and A + C + D (total of 47 genes) were potentially disrupted genes that only exist in control cohort but disappeared in $\textit{PSEN1}$ mutation carrier cohort. Genes from combinations of B + E, B + C + E, and B + C + D + E (total of 13 genes) were potentially emerged risk genes that were differentially correlated with $\textit{PSEN1}$ and/or differentially expressed when comparing $\textit{PSEN1}$ mutation carriers to non-carriers, and they were not correlated with $\textit{PSEN1}$ in control cohort.
Pathway analysis were performed to the disrupted (N = 47) gene groups, emerged (N = 13) gene groups, separately and combined. In the disrupted group, arrhythmogenic right ventricular cardiomyopathy (ARVC) pathway reached significance after multiple testing correction, which contains JUP, ITGB4, and LMNA. Interestingly, a missense mutation (p.As333Gly) in PSEN1 were previously reported in severe progressive dilated cardiomyopathy[170] suggesting a shared common pathway between heart disease and autosomal dominant AD were disrupted in PSEN1 mutation carriers. Among the three ARVC related genes, LMNA is particularly interesting because mutation in this gene alone could result in an extremely rare disease called Hutchinson-Gilford progeria syndrome (1 in 4 million
newborns worldwide). Patients with this disease went through a rapid “premature aging” process with growth failure, fat and hair loss, age-looking skin, and death at average of 14.6 years largely due to cardiac disease or cerebrovascular disease[262]. Although due to very limited autopsy samples, no pathological signs related to dementia or Alzheimer’s disease was identified, this accelerated aging process manifested in this disease is intriguing because aging is the most important risk factor for AD. Previously in another independent study, we also observed a significant increase of LMNA expression in AD brain and significantly associated with plaque load[279]. Notice that Alzheimer’s disease presenilin pathway containing JUP and CD44 was also significant in the disrupted group pathway analysis. In the emerged group, ELMO1 and DOCK1 are related to shigellosis, bacterial invasion of epithelial cells, and integrin signaling pathway (Table 4.4). Interestingly, our group have identified a circular form of DOCK1 (circDOCK1) was significantly up-regulated in ADAD cohort with predominantly PSEN1 mutation carriers. DYNC1LI2 encodes light intermediate chain 2 of dynein protein, which is a family of cytoskeletal motor proteins. It is not surprised to see this gene is related to Huntington Disease pathway. When combing both disrupted and emerged groups, apart from ARVC, shigello also reached significance after multiple testing correction. Shigella, a type of Gram-negative bacteria, could induce shigello after infection with symptoms including diarrhea, fever, and stomach cramps. The infection usually last for 5 to 7 days but there has been evidence suggesting Gram-negative bacterial molecules, such as lipopolysaccharide and E coli K99 pili protein, are associated with AD risk and colocalized with amyloid plaques[286]. Shigella infection related pathway might be involved as an emerged immune response to presenilin pathway disruption in PSEN1 mutation carriers or directly related to sporadic AD susceptibility.
Table 4.4 PSEN1 module disrupted and emerged genes pathway analysis

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4.5 Discussion

To interrogate highly interconnected complex system, network approach not only provides a holistic view of the overall topology but also retains the resolution to the level of each pairwise relationship. With this powerful approach, I analyzed sporadic AD cohort using a system-based approach focusing on the TREM2 module. Using WGCNA, I reconfirmed the observation of MS4A4A and MS4A6A in the TREM2 module, and inferred that MS4A4A might be the key regulator of TREM2 through Bayesian network analysis. This finding replicated previous network analysis that observed the same TREM2-MS4 pattern, and also provided a network explanation of our recent finding about the strong association between MS4A gene cluster and soluble TREM2 level in CSF. I then adapted the approach and applied to ADAD cohorts specially with PSEN1 mutation carriers. With tissues collected from multiple cortical regions in the context of non-demented controls, I constructed a healthy cortical network topology. The network nodes are comprised of genes that are significantly co-expressed with PSEN1 in both and PSEN1 mutation carriers and non-demented controls with no PSEN1 mutation. With functional annotation of the network nodes, I identified 47 genes only present in control cohort that were potentially disrupted in PSEN1 mutation carriers; I also identified 13 genes only present in PSEN1 mutation carriers but not in control cohort that were potentially emerged as downstream transcriptional events acquired into the PSEN1 network. In this list of genes, I highlighted three genes, LMNA, DOCK1, and DYNC1LI2, that are associated to AD in additional studies.

LMNA encodes lamin protein[281], which is important for nucleoskeleton structure in providing mechanical support to nuclear envelope[61, 245]. Apart from the premature aging
syndrome mentioned above in Chapter 4.4.3, other disease phenotypes include muscular dystrophy, lipodystrophy, and cardiomyopathy. In relation to AD, it has been shown that lamin dysfunction has led to neuronal death in tauopathy in *Drosophila*, which is also conserved in human tauopathy[96]. In an interactome study that produced a comprehensive map of molecular interactions derived from yeast two-hybrid and literature curated interactions[42, 182], Alzheimer’s disease and heart disease are close neighbors in the network by sharing several proteins associated with both diseases. Apart from the missense mutation in *PSEN1* that leads to heart disease as I discussed above in Chapter 4.4.3, co-occurrence of cardiovascular disease and AD in elderly suggested additive or synergistic effects on both sides[16].

*DOCK1* encodes a member of the dedicator of cytokinesis protein (Dock) family. *DOCK3* has been found to be associated with AD[149]. This gene’s protein product was originally discovered in a yeast two-hybrid protein-protein interaction screening, which binds to presenilin so it was named as PBP (presenilin-binding protein) in the original paper[149]. We have recently inferred circular RNAs (circRNAs) from RNA-Seq data, which is a type of noncoding RNA that may be involved in AD through a circRNA-mediated “miRNA sponging systems”[175]. Although Dock1 does not bind to presenilin directly[149], circular form of DOCK1 transcript (circDOCK1) was found to be upregulated in both sporadic and autosomal dominant AD. These observations suggested a potential involvement of Dock family in AD pathology through the presenilin pathway.

*DYNC1LI2* encodes light intermediate chain of dynein, which is a motor protein that is required for retrograde axonal transport along microtubules[236, 264]. Dysfunction of this protein leads to disruption of endosomal and lysosomal pathway[257]. Mutations in dynein has
been found in several neurodegenerative diseases suggesting its essential role in neuronal survival, especially for motor neurons[48, 81, 188, 228]. In relation to AD, accumulation of amyloid precursor protein was observed in aged monkey (*Macaca fascicularis*) brains with dynein knockdown[152], and dynein-mediated endocytic dysfunction[256] with increased Rab GTPase level might be involved in this process[152].

With more high-throughput omic data being generated, integrating data from different sources and dimensions will be a promising future direction for network analysis. Apart from multi-dimensional networks, generating hierarchical network (network of networks) could also be an interesting direction. By learning more from the science of complexity, we could gain more insights into the complex systems from gene regulation[123] to human brain[173]. To conclude, my study demonstrated the potential of using both system-based and seed-based network approaches in replicating and discovering AD related genes and their interactions. In sporadic AD cohort, I identified *MS4A4A* might be a key regulator for *TREM2*. In autosomal dominant AD cohort, I identified total of 60 genes that are disrupted or emerged as a consequence of *PSEN1* mutation, in which I highlighted three genes with intriguing links to AD. Many of the remaining genes that are not discussed in detail in this chapter are also found to be associated with AD, such as *ABCA2*[176], *GFAP*[147], *CXorf36*[207], suggesting a great potential of using network analysis to generate working hypothesis.
Chapter 5: Conclusions and future directions
5.1 Dissertation work contributed to AD research

In Chapter 2, a deconvolution pipeline for bulk RNA-Seq was developed to account for cell type specific effects in brain tissues. Due to disease pathology, cell type balance is disrupted in Alzheimer’s Disease (AD) brain, which is a key feature in neurodegeneration that has often been overlooked in transcriptome research. With deconvolution methods to better delineate cell population changes in disease condition, it would help interpret results and reveal transcriptional changes in a cell type specific manner.

In Chapter 3, using cell type proportion as quantitative trait, a common pathway underlying aging brains has been identified in the presence or absence of neurodegenerative disease symptoms. A protective variant of TMEM106B, which was previously identified with a protective effect in FTD, was identified to be associated with neuronal proportion in aging brains, suggesting a common pathway underlying neuronal protection and cognitive reservation in elderly. This extended analysis yield from deconvolution results from Chapter 2 demonstrated one promising application of deconvolution followed by cell type QTL analysis in identifying new genes or pathways underlying neurodegeneration or aging in general.

In Chapter 4, using network analysis I replicated and reconfirmed the co-expression pattern between MS4A gene cluster and TREM2 in sporadic AD, from which further evidence was inferred from Bayesian network analysis to show that MS4A4A might be a potential regulator of TREM2 that is validated by in-vitro experiments. In Autosomal Dominant AD (ADAD) cohort, disrupted and acquired genes were identified from PSEN1 mutation carriers. Among the genes, previous identified AD related gene and pathways were revealed together with
novel findings. These results demonstrated the great potential of applying network approach in identifying disease associated genes and the interactions among them.

5.2 Aging, proteinopathy, and neurodegeneration

Apart from clinical criteria, diagnosis of neurodegenerative disorders heavily hinges on proteinopathies observed either from longitudinal brain imaging and Cerebrospinal fluid data and postmortem neuropathology. Although each disease has its own characteristic protein(s), pathological proteins from a different neurodegenerative disorder are also observed in patient autopsies. For example, AD is characterized by amyloid β and tauopathy; Parkinson’s Disease is characterized by alpha-synuclein; FTD is characterized by TDP-43 pathology. However, in AD post-mortem tissues alpha-synuclein and TDP-43 are often detected in. Similarly, amyloid β and tau are also present in PD. TDP-43 is not only identified in FTD brains, but it also presents in various neurodegenerative disorders, such as AD, ALS, hippocampal sclerosis as I have discussed in Chapter 3.5. Based on what I have learnt from the field and observations obtained from this dissertation work, a model is summarized and depicted in Figure 5.1.

Initially, soluble protein substrates in their normal folding states are performing normal functions in the brains. However, due to triggering-events that could be genetic, lifestyle or environmental factors, such as microbial infection, traumatic brain injury, toxic metal exposure, and even lack of sleep, the normal protein substrates become insoluble through a mis-folding process and start to aggregate. Regarding the initial formation of protein aggregates, a seeding hypothesis has been proposed[139]. Taking amyloid formation as an example, an initial nucleation event generates a seed that later initiates the following pathogenic accumulation of amyloid β proteins in an exponential way of aggregation. Then this pathogenic aggregation starts
to propagate based on well-established prion observations that a diseased protein could convert a normal protein into a diseased state so that this process is able to propagate to other unaffected proteins. One study also demonstrated that the proteins with prion-like characteristics are also transmissible from patient to patient through injection of cadaveric pituitary-derived growth hormone[140] that induced AD pathology in patients without genetic risk factors.

What type of mis-folded proteins are generated depends on the genetic background of the individual. For example, if one person carries mutations or risk variants in AD risk genes, such as \( \text{APP, PSEN1, PSEN2, APOE, TREM2} \), this individual will be susceptible to amyloid \( \beta \) and tau aggregation. Carriers of PD risk variants, in genes such as \( \text{PARK2, LRR2, PINK1, SNCA} \), will be susceptible to alpha-synuclein aggregation. And similarly, carriers of variants in FTD risk genes, such as \( \text{MAPT, GRN or C9orf72} \), will be susceptible to tau and TDP-43 aggregation. As it was discussed above, different categories of proteinopathies are not mutually exclusive in their distribution, which means alpha-synuclein protein aggregation could be observed in AD patients with amyloid \( \beta \) and tau pathologies. Thus, a patient as depicted in Figure 5.1 is clinically diagnosed with AD, but this individual also harbors multiple other proteinopathies that have not reached the threshold for clinical manifestation. This observation could be explained by an omnigenic model that apart from core pathways of AD risk genes, a collection of low-effect risk genes for other neurodegenerative disorders contribute non-trivial effects to the pathologic protein aggregation process. Apart from risk factors, there are also protective factors that could help ameliorate the damaging effect brought by toxic proteins that protect individuals with protein aggregation pathology but no cognitive deficits.
Figure 5.1 Proteinopathy model in neurodegenerative disorders. Triggered by various factors such as familial mutation, microbial infection, traumatic brain injury, toxic metal exposure, or general aging process, normal proteins may become different types of misfolded proteins based on a subject’s specific genetic or epigenetic architecture as shown in A) that form alpha-synuclein, amyloid beta, or TDP-43 proteinopathies that belong to different neurodegenerative disease categories. B) Accumulated proteinopathies may lead to neuronal loss. C) As disease progresses, the subject may experience cognitive deficit with more accumulated proteinopathies. D) A patient diagnosed with AD may have alpha-synuclein and TDP-43 deposition apart from amyloid plaques that suggests neurodegeneration shall be considered as a continuous trait rather than distinctive disease categories.
5.3 Future directions in developing therapies for AD

A prevalence study focusing on preclinical and clinical AD prevalence showed that there are eight times more people in pre-symptomatic phase than people in clinical phase[39]. People with preclinical AD have either amyloidosis or neurodegeneration or both, but have no clinical manifestation of symptoms. This data showed great promise for preventive therapies that by either slowing down disease progression or delaying onset, it will be possible to protect a large number of people from developing AD during their lifetime. Currently, more than half of the disease modifying therapies are focusing on anti-amyloid approach, including immunotherapy, BASE inhibitor, and anti-aggregation. However, there have no signs of any success so far in this path. One problem with some clinical trial designs are the patients recruited for the trials are usually in their middle or late stage of AD. As it has been shown above, during the long preclinical phase, especially in the sporadic form of neurodegenerative disorders, neuropathological and neurodegenerative changes have occurred long before any clinical symptoms. However, by the time of clinical manifestation, it is usually too late for any treatment due to the extensive brain damage. Clearance of aggregated proteins may not be effective to compensate cognitive deficit due to massive neuronal loss. If neuronal loss is not reversible in the current situation, early intervention, amyloid clearance or trigger-targeting therapies should be performed earlier during disease progression to target people still in their pre-symptomatic phases. Apart from the amyloid pathway, other potential pathways to target are the lysosomal and autophagy pathway to facilitate toxic protein degradation. In addition, targeting immune pathway could prevent vicious cycles of proinflammatory responses and boost immune resilience to infections. All these pre-clinical therapies heavily rely on achieving early diagnosis in the general population. To advance early diagnosis, the development of non-invasive and accurate
diagnosis tools is highly demanded to predict disease at early stages before symptom onset. We also need to advance our understanding of genetic and environmental triggers of AD to identify and better target susceptible cohorts.

Given the complexity of neurodegenerative disorders, we also need to tailor therapies differently for people with different genetic backgrounds. This idea of precision medicine has gained success in other medical fields, for example cancer treatment, from which we can learn. With better understanding of AD etiology, test results shall be interpreted in the context of person specific genetic architecture. Studies with more diverse cohorts shall be supported to understand ethnic and sex differences, which should also be considered in evaluating drug responses, dosages and side effects during clinical trials. Last but not least, cognitive or motor function improvement therapies and health care facilities specially designed for people with dementia or motor deficits are also needed for patients with neurodegenerative disorders. Regarding recent interests in finding protective factors, neuronal protection therapies could prevent the major detrimental consequence. As documented in this dissertation work, using cell type compositions inferred from deconvolution as disease endophenotypes, I identified protective variants in \textit{TMEM106B} gene that may have neuronal protection effect in general aging groups independent of disease status, which could help understand the relationship between aging and neuronal survival and be a potential target for neuronal protection therapies.
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Washington University – St. Louis, MO, USA  
Doctor of Philosophy, May 2019  
Major in Neuroscience  
Cognitive, Computational & Systems Neuroscience Curriculum Pathway

Purdue University – West Lafayette, IN, USA  
Bachelor of Science, May 2010  
Major in Biology

China Agricultural University – Beijing, China  
Bachelor of Science, May 2010  
Major in Biological Science

**Research Experience**

Carlos Cruchaga Lab, Washington University, St. Louis, MO  
Sept 2015 – Present

Project: Integrative Analysis to Investigate Complex Interaction in Alzheimer’s Disease

- Developed a semi-supervised NMF deconvolution pipeline to resolve tissue heterogeneity in bulk RNA-seq.
- Analyzed genetic variants and transcriptome data in Alzheimer's disease to better understand disease etiology and to identify potential therapeutic targets.
• Applied machine learning algorithms and graph-based network analysis to next generation sequencing and proteome data to identify risk factors associated with Alzheimer’s Disease.
• Collaborated on multi-omic projects including CSF and plasma protein, cell free RNA, circular RNA, single nuclei RNA sequencing data analysis.

John Pruett Lab, Washington University, St. Louis, MO Nov 2012 – Aug 2014

Project: Exploring Cortical Biomarkers for ASD with fcMRI Functional Parcellation

• Applied community detection and image gradient detection algorithms to analyze infant brain imaging data to identify functional areal differences in infants and adults and discover pre-clinical biomarkers for Autism disorders in infants.

Yuk Fai Leung Lab, Purdue University, West Lafayette, IN Jan 2009 – May 2011

Project: The Role of Phenylthiourea in Zebrafish Eye Growth Regulation

• Measured the smaller eyes caused by phenylthiourea (PTU) treatment.
• Proposed and tested the thyroid hormone hypothesis that the smaller eye in the PTU-treated larvae may be caused by a lower thyroid activity.
• Published one research article and one review article as first authors during undergraduate research.
• Received two internal research awards and one external research grant during undergraduate research.

De Ye Lab, China Agricultural University, Beijing, China Mar 2007 – Jan 2008

Project: KD616 Promoter of Arabidopsis Thaliana Clone, Localization, and Expression Analyses

• Constructed a bacterial plasmid of predicted KD616 promoter fused with GUS reporter gene, infected Arabidopsis Thaliana, selected and stained the seeds of transgenic plants to observe GUS signals.

Employment

Research Assistant Aug 2010 – May 2011

Yuk Fai Leung Lab, Purdue University, West Lafayette, IN

• Conduct research to elucidate the role of phenylthiourea in zebrafish eye growth regulation.
• Conduct routine fish room and lab maintenance.
• Assisted in performing immunostaining, quantitative PCR and statistical analyses using R for several ongoing projects.
Department of Biological Sciences, Purdue University, West Lafayette, IN

- Tutored freshmen and sophomores in introductory level biology courses.

**Selected Publications**

- **Li Zeran**, Farias FG, Dube U, et al. The *TMEM106B* rs1990621 protective variant is also associated with increased neuronal proportion. bioRxiv 2019.

**Academic Achievements and Awards**

- Boeing Patent Challenge Second Prize Dec 2015
- Graduated with Distinction/Honor Curriculum May 2010
- Dr. William H. Phillips Undergraduate Research Internship Mar 2010
- Sigma Xi Grants-in-Aid for Vision-related Research Jan 2010
- Sandy and Zippy Ostroy Research Experience for Undergraduates Award Dec 2009
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