1 Molecular estimation of neurodegeneration pseudotime in older brains

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26 Abstract

27	Therapeutic treatments for late-onset Alzheimer's disease (LOAD) are hindered by an incomplete
28	understanding of the temporal molecular changes that lead to disease onset and progression. Here, we
29	evaluate the ability of manifold learning to develop a molecular model for the unobserved temporal
30	disease progression from RNA-Seq data collected from human postmortem brain samples collected
31	within the ROS/MAP and Mayo Clinic RNA-Seq studies of the AMP-AD consortium. This approach
32	defines a cross-sectional ordering across samples based on their relative similarity in RNA-Seq profiles
33	and uses this information to define an estimate of molecular disease stage - or disease progression
34	pseudotime - for each sample. This transcriptional estimate of disease progression is strongly concordant
35	with burden of tau pathology (Braak score, $P = 1.0 \times 10^{-5}$), amyloid pathology (CERAD score, $P = 1.8 \times 10^{-5}$)
36	⁵), and cognitive diagnosis ($P = 3.5 \times 10^{-7}$) of LOAD. Further, the disease progression estimate
37	recapitulates known changes in cell type abundance and impact of genes that harbor known AD risk loci.
38	Samples estimated to reside early in disease progression were enriched for control and early stage AD
39	cases, and demonstrated changes in basic cellular functions. Samples estimated to reside late in disease
40	progression were enriched for late-stage AD cases, and demonstrated changes in known disease processes
41	including neuroinflammation and amyloid pathology. We also identified a set of control samples with
42	late-stage estimated disease progression who also showed compensatory changes in genes involved in
43	affected pathways are protein trafficking, splicing, regulation of apoptosis, and prevention of amyloid
44	cleavage. In summary, we present a disease specific method for ordering patients based on their LOAD
45	disease progression from CNS transcriptomic data.

46 Introduction

47 Late-onset Alzheimer's disease (LOAD) is a devastating illness with no effective disease modifying

48 therapy, owing to a 99.6% failure of clinical trials¹. There is a growing consensus that the most effective

49 treatments will intervene early in disease progression and halt disease pathophysiological processes prior

50 to conversion to $LOAD^2$. In addition, there is increasing recognition that LOAD may in fact be a

spectrum of related diseases that have similar clinical and neuropathological manifestations^{3,4}. Devising
successful therapeutic strategies will likely require targeting potentially diverse early-stage disease
processes that occur prior to a high burden of neuropathology or cognitive impairment.

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55 Current approaches to identify AD affected individuals include *in vivo* measures of the pathological 56 hallmarks of disease – amyloid, tau, and neurodegeneration – via CSF biomarkers for amyloid and tau⁵, 57 positron emission tomography for amyloid and tau (PET)⁶, and structural and functional MRI of 58 neurodegeneration. Cognitive assessments are used to estimate disease burden⁷, although measurable 59 cognitive impairment generally indicates a sustained burden of neuropathology and advanced 60 neurodegeneration. Based on biomarker studies of AD, by the time cognitive decline becomes detectable, 61 neuropathological changes of AD have already occurred, first in Aß and subsequently in tau related 62 measures⁸ and therefore cannot be used to select patients for early disease stage studies. Furthermore, 63 while these measures of disease progression capture the overall increase in burden of pathology and 64 cognitive decline, they do not necessarily identify the dysfunctional molecular mechanisms that lead to 65 neuropathology and cognitive decline. There are likely many independent patient specific molecular pathways present at an early stage in disease that then contribute to later stage disease progression 9,10 . 66 This motivates the need to identify these early stage molecular mechanisms driving disease progression. 67 68 69 The Accelerating Medicines Project for Alzheimer's Disease (AMP-AD) consortia have generated 70 genome-wide transcriptomics of post-mortem brain tissue from patients across a broad range of

Alzheimer's disease neuropathological progression – including individuals with various stages of AD
neuropathology and those who lack AD neuropathology, but who may in face harbor early stage disease
molecular processes. We therefore sought to chart the molecular progression of the disease as reflected in
the aggregate behavior of the CNS transcriptome across these individuals. While standard approaches
such as differential expression or coexpression analyses have proven informative^{11–15}, these analyses do

not infer the relative stage of disease progression or identify distinct disease subtypes. Here we propose

an approach to analyze population level RNA-seq data from post-mortem brain tissue to learn a tree
structured progression (Figure 1) that represents distinct sub-types of disease and the relative progression
of disease across patients. With this approach, we identify potentially generalizable trajectories of LOAD
across heterogeneous patient populations at all stages of disease. Furthermore, we characterize molecular
pathways that define disease stages – a potential source of new biomarkers and therapeutic interventions
for early-stage disease processes along multiple different disease trajectories.

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84 To learn the molecular disease staging and neuropathologic progression tree we use a manifold learning method¹⁶. Manifold learning refers to a group of algorithms that aim to recover the low dimensional 85 86 subspace underlying a high dimensional dataset. Previous authors use manifold learning to estimate 87 disease progression from neuroimaging data¹⁷ and to study lineage commitment of cells during differentiation from single cell RNA-seq (scRNA-seq)¹⁸⁻²¹. To our knowledge, manifold learning has not 88 89 been used to estimate disease progression and/or disease stages from bulk RNA-seq data derived from 90 post-mortem tissue. Henceforth, we refer to manifold learning, lineage inference interchangeably in 91 reference to the construction of the inference of a neuropathologic progression tree. We demonstrate that 92 these tools can estimate the disease staging and progression tree (Figure 2) from bulk RNA-Seq data 93 collected from post-mortem brain tissues in a case/control cohort. Moreover, these trees show clear 94 LOAD staging, enable the study of cell type specific effects of LOAD, and allow the identification of 95 genetic factors driving disease progression.

96 **Results**

97 Unsupervised manifold learning distinguishes pathologically defined LOAD from control

98 We first quantify the bulk RNA-Seq data from the ROS/MAP and Mayo Clinic cohorts into gene counts

99 and remove any batch effects introduced due to sequencing runs using standard count normalization (see

100 Methods). The data from the ROS/MAP cohort is sampled from the dorsolateral prefrontal cortex

101 (DLPFC), and the data from the Mayo Clinic cohort is sampled from the temporal cortex (TCX). The full pipeline we use for RNA-Seq data generation and quality control was recently reported²². The entire 102 103 transcriptome comprises many genes which do not have measurable expression or vary across case/control samples, which we remove in order to reduce the noise in manifold learning¹⁹. To do this, we 104 105 first perform differential expression analysis between case/control samples separately for each study and 106 retain genes that reach an FDR of 0.10. To test if this biased the disease lineage inference, we also 107 perform manifold learning using only genes with high variance across samples, and we see a strong 108 concordance with disease lineages inferred with differentially expressed genes (Figure S4). We infer the 109 disease lineage for each brain region on this subset of retained genes (Figure 2A-B). Furthermore, we 110 observe strong evidence of sex heterogeneity when performing the manifold learning approach, and find 111 that the manifolds inferred for female only samples are much more robust than for male samples. This matches previous observations concerning disease specific sex heterogeneity 22 . We only show results for 112 113 manifolds inferred on female samples.

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115 We first visualize the clinical diagnosis of the samples on the inferred disease staging tree to verify that 116 there is indeed separation of AD patients across the tree. To determine if inferred tree structure is an 117 accurate model of disease progression, we introduce the notion of disease pseudotime which is the 118 geodesic distance along the tree from an inferred initial point to the point of interest as a quantitative 119 linear measure of LOAD stage. We scale this estimated disease pseudotime to lie in the range [0,1] to 120 make the effects comparable between the two studies (and brain regions). We show that for LOAD cases compared to controls there is a significant association (P = 0.02 in Mayo and $P = 2.0 \times 10^{-6}$ in ROS/MAP. 121 122 logistic regression) between the estimated pseudotime and AD case/control status (Figure 2C).

123

We test whether genes in loci that have been implicated in genome wide association studies of LOAD are associated with inferred disease pseudotime. We use the highest ranked LOAD GWAS genes (60 genes in total)²³, **Table S1**, and compute the correlation between their expression and inferred pseudotime (Figure 2D). When compared to the background of all genes, we see that there is a significant increase in
positive correlation with disease pseudotime for implicated LOAD GWAS genes (P-value: 7.3x10⁻⁵ in
Mayo and 5.6x10⁻³ in ROS/MAP). Furthermore, this does not appear to be driven by a small subset of
outlier genes, but by the majority of the distribution of LOAD GWAS genes. The fact that AD GWAS
loci genes have expression associations with pseudotime likely implies that the AD risk variants at these
are also eQTL as previously shown²⁴⁻²⁷ and/or are members of co-expression networks that are
differentially expressed in AD^{13,28}.

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135 To further explore the relationship between inferred disease stage and LOAD, we test for its association 136 with neuropathological and clinical measures of LOAD severity, namely: i) Braak score, ii) CERAD 137 score, and iii) cognitive diagnosis. The ROSMAP study has numeric scores for these categories available 138 as covariates for each sample. Braak is a semi-quantitative measure that increases with tau pathology 29 139 and CERAD is a semi-quantitative measure of density of neuritic plaques³⁰. We overlay these scores on 140 the inferred manifold for the DLPFC brain region (Figure 3A). We observe a progressive increase in tau, 141 amyloid, and cognitive burden as we traverse the inferred disease manifold (Figure 3A). This is further 142 quantified by characterizing the relationship between branches of the inferred manifold and Braak, 143 CERAD, and cognitive diagnosis (Figure 3B). We observe significant associations between pseudotime and Braak score (P=1.0x10⁻⁵), CERAD score (P=1.8x10⁻⁵), and cognitive diagnosis (P=3.5x10⁻⁷). 144

145 Inferred staging recapitulates known biology of AD

To demonstrate that the inferred disease pseudotime recapitulates known biology of LOAD, we test for association between inferred disease stage and both the cellular response to disease and the genetics of the disease. A prominent hypothesis in AD is that the effects of the disease vary across different brain cell types, specifically neurons and glial subtypes. Current understanding of the cell biology of the disease implicates progressive neuronal loss and increase in gliosis³¹. To test if the inferred pseudotime aligns with existing cell type specific hypotheses regarding AD, we first selected from the genes used in lineage

152	construction the marker genes for four key cell types: neurons, astrocytes, microglia, and
153	oligodendrocytes based on a previously published brain cell atlas ³² (Table S2). We then calculate the
154	normalized mean expression for the marker genes of each cell type and fitted a linear model to the mean
155	expression with disease pseudotime as the dependent variable. We find that, in both studies, the cell
156	specific marker gene levels show a statistically significant linear dependence on pseudotime (Table S3).
157	Fitted effects recapitulate known neuropathologic changes which occur in AD, namely: i) a reduction in
158	the neuronal populations as AD progresses, and ii) an increase in expression associated with activation of
159	microglia, astrocytes, and oligodendrocytes as AD progresses (Figure S5).
160	
161	Next, we test for association between assigned lineage state in ROS/MAP (DLPFC) and Mayo (TCX) and
162	APOE e4 status (Figure S6). For reference, the inferred trees for TCX and DLPFC each resolve into 6
163	branches (Figure 4A,S7). Carriers of the APOE e4 allele are significantly enriched on the State 4 branch
164	in TCX (P-value = 0.027, unadjusted), and suggestively enriched on the State 5 branch (P-value = 0.06,
165	unadjusted), compared to the State 1 branch (logistic regression).

166

167 *Genetic factors associated with inferred disease staging*

168 Lineage inference of LOAD transcriptomes provides a quantitative measure of disease progression for 169 genetic associate testing, and the significantly greater correlation between pseudotime and gene 170 expression for known LOAD risk genes (Figure 2D) suggests that the observed differences in disease 171 trajectories are influenced by genetic factors. To test this hypothesis, we perform single variant analysis 172 using whole-genome sequencing data for 305 patients from the ROS/MAP and 131 patients from the 173 Mayo cohort. Despite the limited sample size, resulting in lack of statistical power to discover genome-174 wide significant associations, multiple variants reach a genome-wide suggestive threshold of $p < 1 \times 10^{-5}$ 175 (Table S4). We do not see evidence of population stratification in the analysis (Figure S8-S9). Notably, 176 the most significant association with pseudotime for the ROS/MAP cohort is observed at the PTPRD 177 locus (rs7870388, p = 1.31×10^{-6}) (Figure S10, Table S4). The *PTRPD* locus is associated with the

susceptibility to neurofibrillary tangle independent of amyloid deposition in the ROS/MAP cohort³³. For 178 the Mayo Clinic cohort, known LOAD variants in the APOE (rs6857, $p = 9.18 \times 10^{-6}$) and BIN1 179 180 $(rs62158731, p=4.68 \times 10^{-5})$ loci overlap with variants associated with inferred disease stage (Figure **S10. Table S4**)³⁴. When comparing our association results for inferred disease stage with summary 181 182 statistics from a large-scale case-control approach, we identify multiple variants which have been 183 previously associated with LOAD in the IGAP cohort (Table S5). Furthermore, we identify several 184 potential novel candidate genes associated with inferred disease stage (ADAMTS14, IL7, MAN2B1) linked 185 to immune and lysosomal storage function (Figure S10, Table S4). IL-7 has been proposed as an inflammatory biomarker for LOAD that correlates with disease outcome and severity³⁵. ADAMTS14 is 186 187 part of a locus that has been previously linked with Alzheimer susceptibility and plays an important role

- 188 in the regulation of immune function via TGF-beta signaling.
- 189 New disease insights identified from inferred disease lineages

Another important direction of study in the field of Alzheimer's is the identification of disease subtypes, which has so far predominantly been done using imaging data³⁶. The branches of the inferred disease trees provide a new transcriptomic-based approach to identify disease subtypes. In both brain regions and in two separate cohorts, there were two distinct early-lineage branches corresponding to predominantly control samples, which we interpret as different initial paths towards the disease. Similarly, both brain regions feature several distinct branches with predominantly LOAD samples (Figure 2A-B).

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Branch-specific differential expression patterns. To study the genes and pathways specific to each branch,
we perform a branch-specific differential expression analysis with an ANOVA model using the branches
with the highest proportion of controls as the reference branch for DLPFC (**Table S6**) and TCX (**Table**S7). We see many genes are differentially expressed between the control branch and branches that are
enriched in the affected individuals (**Table S8**). Next, we performed an enrichment analysis on each of
these differentially expressed gene sets with the enrich R³⁷ package for Gene Ontology³⁸ annotations

(Methods). The results of this enrichment analysis for DLPFC and TCX tissues (Table S9-S10). Only
gene sets with significant enrichment are shown (FDR adjusted p-value < 0.05). Overall, we see a pattern
of loss of expression of basic cell biology mechanisms in early-stage branches including RNA splicing,
mitochondria function, protein transport, and DNA repair. Late-stage branches were characterized by
increased immune response (e.g. TGFb/WNT signaling) and apoptotic activity (Table 1).

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209 While studying the different branches in the two brain regions, we observe a branch (branch 5) that 210 corresponds to a group of predominantly neuropathological control samples from the Mayo RNA-seq 211 cohort that were in close proximity to a branch with predominantly LOAD samples (branch 4) on the 212 inferred disease lineage (Figure 4A). However, most of the samples on branch 5 are neuropathological 213 controls as defined by the Mayo diagnostic criteria. We bi-cluster the mean expression of genes in each 214 branch and the branches themselves (**Methods**). This clustering analysis (**Figure 4B**) shows that the 215 closest branch to this potentially disease resistant branch contains the highest proportion of AD samples. 216 While the stage proximity implies some transcriptomic similarity between these controls and nearby 217 cases, we also see a secondary cluster of genes with increased expression in the resistant state while 218 having reduced expression in all other states. We perform an enrichment analysis on this set of genes and 219 find significant GO terms corresponding to: protein transport (GO:0015031), regulation of mRNA 220 splicing, via spliceosome (GO:0048024), negative regulation of apoptotic process (GO:0043066), and 221 regulation of amyloid-beta clearance (GO:1900221) (Cluster4, Table S11). It is possible that these 222 potentially disease resistant individuals have compensatory mechanisms which suppress the hallmarks of 223 disease despite sharing gene expression patterns with pathologically affected individuals. These 224 observations are preliminary, and would need to be replicated in a second cohort to verify the existence of 225 a disease resistant gene expression signature.

226 Discussion

227 Here we proposed a novel approach to infer the Alzheimer's disease severity and disease subtypes in an 228 unsupervised manner from post-mortem bulk RNA-seq data that gets directly at the challenge of 229 identifying the temporal progression of disease in the disease resident tissue. Our strategy utilized a 230 manifold learning approach to infer a disease progression tree from cross-sectionally collected patient 231 samples from two different brain regions. The underlying assumption of our approach is that the inferred 232 disease progression from cross sectional samples serves as a proxy for the unobserved progression of the 233 disease across subtypes of LOAD. We validated this hypothesis through comparisons with 234 neuropathological measures of disease stage severity and against known cell type specific effects caused 235 by the disease. Furthermore, this approach provides clues to better understanding the molecular 236 heterogeneity of disease by identifying specific pathways that are dysregulated in subsets of patients at 237 different disease stages. This opens up the possibility of better patient stratification and precision 238 medicine.

239

240 We observed that different biological processes vary as a function of inferred disease stage, and that 241 early-stage disease processes potentially include RNA-splicing, mitochondrial function, and protein 242 transport – implicating multiple basic cell biology mechanisms as potential early stage disease processes 243 for further study in relevant model systems. Additionally, the manifold learning method identified 6 244 potential subtypes of LOAD from RNA-seq (i.e. branches) suggesting the LOAD populations should be 245 stratified by better biomarkers with tailored treatment strategies. To identify and test these stratifications 246 future studies should focus on longitudinal cohorts of patients with rich molecular and imaging data to be 247 able to identify biomarkers that can accurately and precisely stratify patients into the underlying 248 molecular subtypes in terms of the molecular characteristics of their transcriptome and different relative 249 stages of disease. Furthermore, we observe a potential disease resistant sub-type of patients. This disease 250 resistance should be tested in disease model systems, to identify if neuropathological readouts can be

modified by altering the function of the pathways identified in our analysis (e.g. APP processing, RNA
splicing, apoptosis, protein trafficking). While this preliminary observation needs to be validated in
another cohort, it has the potential to be a novel source of hypotheses concerning new therapeutic
development. Specifically, for constructing better combination therapy hypotheses that may confer
neuroprotection, even in patients that are mildly affected by disease.

256

257 LOAD is a complex and heterogeneous disease encompassing a broad spectrum of clinical symptoms. 258 Disease progression can vary widely between patients leading to different rates of cognitive decline. 259 Several lines of evidence suggest that these differences in progression are modified by multiple genetic factors affecting the transition from one pathological state to another^{39,40}. However, it has remained 260 261 difficult to assess the role of genetic variants affecting disease trajectories by case-control approaches 262 alone. Here, we showed that our novel expression trait pseudotime might be used as a molecular 263 phenotype to identify known and novel AD loci associated with different disease progression states across 264 AD patients. Despite a limited sample size, we identified previously associated AD candidate loci in the 265 ROSMAP (*PTPRD*) and Mayo (*BIN1*, *APOE*) cohorts with suggestive significance ($p < 1 \times 10^{-5}$). 266 Variants in *PTPRPD* have been associated with the susceptibility to neurofibrillary tangles, independent 267 of amyloid burden. This is in line with the results from the differential gene expression analysis of 268 pseudotime branches showing an enrichment of molecular pathways implicated in TAU pathology. 269 Furthermore, our analysis revealed several novel loci linked to immune function (ADAMTS14, IL7) and 270 neurotransmitter signaling (CHRM2, CHRM3) processes associated with disease pseudotime (Table S4). 271 Future studies will be needed to replicate these findings in independent cohorts of LOAD and validate the 272 role of candidate genes in LOAD related disease progression by first identifying peripheral biomarkers 273 that correspond to this molecular definition of disease stage, and then testing for GWAS association with 274 that disease stage. Subsequent results can improve functional interpretation by linking candidate genes 275 with ordered pathological processes.

277 Methods

278 RNA sequencing

- 279 The details of the sample collections, postmortem sample characteristics, the tissue and RNA
- 280 preparations, the library preparations and sequencing technology and parameters, and sample quality
- 281 control filters are provided in previously published work^{41,42}. Furthermore, details of the bioinformatic
- pipeline used to generate count level data has been previously described²². Briefly, reads were aligned to
- the GENCODE24 (GRCh38) reference genome with STAR⁴³, and gene counts generated using the
- HTSeq algorithm⁴⁴. Genes that had more than one counts per million total reads total reads in at least
- 285 50% of samples in each tissue and diagnosis category were used for further analysis.
- 286

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287 Differential Expression analysis on Mayo and ROS/MAP cohorts

288 For gene filtering we used false discovery rate of 0.05 from the previously published differential

expression analysis of Mayo and ROS/MAP RNA seq data²². Briefly, case control status was harmonized
 across the Mayo and ROS/MAP cohorts, where controls were defined as individuals with a low burden of

amyloid and tau based on CERAD and Braak scores, and cases with a high burden. Furthermore in

251 anyloid and tau based on CERAD and Braak scores, and cases with a nigh burden. Furthermore in

293 cases have probably AD^{22} . Differential expression analysis was run on suitably normalized data – using

ROS/MAP, clinical diagnosis was also used with controls having to have no cognitive impairment, and

- conditional quantile normalization to account for variation in gene length and GC content, removing
- sample outliers, covariate identification adjustment, with sampling abundance confidence estimated using
- a weighted linear model with the voom-limma package 22,45,46 . A fixed/mixed effect linear model is used
- 297 to fit the differential expression model on the normalized data²².

298

299 Manifold learning for LOAD

300 Manifold learning refers to a group of machine learning algorithms that recover a low dimensional
 301 subspace underlying a high dimensional dataset. Manifold learning approaches are typically used in

302 datasets or applications where data samples lie on an underlying low dimensional latent space (e.g. a tree, 303 a line, a curved plane). The low dimensional space is learned via a projection from the high dimensional 304 space of the observed data (e.g. RNA-seq profiles across hundreds of patient samples) down to a low 305 dimensional space with suitable regularization constraints to enforce smoothness and the structural 306 constraints of the low dimensional space. (Figure 1A). Due to the necessary assumption of an underlying 307 latent subspace, manifold learning is commonly used in applications where it is known that the observed 308 data is obtained from a progression of some kind; e.g., i) to infer the temporal ordering of a sequence of 309 images, or ii) to infer the approximate lineage of cells in a differentiation trajectory using single cell 310 RNA-Seq data (Figure 1B-C). 311 312 Here, we repurpose methods originally developed for learning cell lineage using scRNA-Seq data, to infer 313 the staging of Alzheimer's disease (AD) using bulk RNA-Seq data from post-mortem brain samples with 314 known AD diagnosis status. Since bulk RNA-Seq has many of the same sampling and distributional 315 properties as scRNA-Seq, we observe that scRNA-Seq methods are applicable with no additional 316 modifications. As such, we use the DDRTree manifold learning approach available in the Monocle 2 R 317 package ¹⁹. However, we also show that the estimated staging of disease is quite similar across some of 318 the other common methods used for scRNA-Seq lineage estimation (Figures S1-S3). 319 320 The RNA-Seq data used in this study was generated from post-mortem brain homogenate samples, and 321 obtained from two separate studies that are a part of the Accelerating Medicines Partnership in 322 Alzheimer's Disease (AMP-AD) consortium, namely: i) the Religious Orders Study and the Memory and Aging Project (ROSMAP)^{47,48}, and ii) the Mayo RNA-seq study⁴⁹. For this paper, we focused our 323 324 analysis on the temporal cortex (TCX) and dorsolateral prefrontal cortex (DLPFC) tissue samples. Within 325 the Mayo RNA-seq study the TCX samples are derived from individuals neuropathologically defined as

either aged controls, LOAD cases, Progressive Supranuclear Palsy (PSP) cases, or pathological aging

327 (PA) cases ⁴⁹. The ROSMAP study is a prospective longitudinal cohort of an aging population, and has

samples from participants with clinical and neuropathological diagnoses of LOAD⁴², aged controls, and
 individuals with mild cognitive impairment. Furthermore, the results presented in the main paper are
 from female samples only, as we observed significant sex differences in the transcriptomic data consistent
 with current knowledge of sex differences in LOAD^{50,51}, making a common analysis of both sexes
 untenable.

333

334 *Manifold learning using Discriminative Dimensionality Reduction Tree (DDRTree)*

335 DDRTree is a manifold learning algorithm that infers a smooth low dimensional manifold by an approach 336 called reverse graph embedding. Briefly, the algorithm simultaneously learns a non-linear projection to a 337 latent space where the points lie on a spanning tree. A reverse embedding is also simultaneously learned 338 from the latent space to the high dimensional data. Mathematically, the DDRTree algorithm can be posed 339 as the following optimization problem:

$$\min_{W,Z,B,Y,R} \sum_{i=1}^{N} ||x_i - Wz_i||^2 + \frac{\lambda}{2} \sum_{k,k'} b_{k,k'} ||Wy_k - Wy_{k'}||^2 + \gamma \left[\sum_{k=1}^{K} \sum_{i=1}^{N} r_{i,k} ||z_i - y_k||^2 + \sigma r_{i,k} \log(r_{i,k}) \right]$$

s.t. B represents a spanning tree,

$$W^T W = I, r_{i,k} \ge 0, \sum_{k=1}^{K} r_{i,k} = 1$$

340

Here, $\{z_i\}_{i=1}^N \in \mathbb{R}^{genes}$ represents RNA-Seq data from each patient sample, $\{z_i\}_{i=1}^N \in \mathbb{R}^2$ represents the latent representation of each sample as inferred by the algorithm, $\{y_k\}_{k=1}^K$ represents the centers of clusters in the dataset, $W \in \mathbb{R}^{2 \times genes}$ represents an inverse mapping from the latent space to the high dimensional space of RNA-Seq data, $B \in \mathbb{R}^{K \times K}$ represents a spanning tree on which the centers of the clusters lie and $R \in \mathbb{R}^{N \times K}$ captures the soft clustering information of samples in the dataset. The first term of the optimization problem is responsible for learning a low dimensional representation of the data such that an inverse mapping exists to the high dimensional data points, the second term learns the tree

348	structure of the points and the third term learns a soft clustering for the latent dimension points as well as
349	the centers of the clusters. Despite the non-convexity of the problem, each individual optimization
350	variable can be solved for efficiently using alternative minimization as described previously ⁵² . This
351	algorithm was implemented using the Monocle package in R ¹⁹ . The code to infer the lineage in Mayo is
352	available here <u>https://github.com/Sage-</u>
353	Bionetworks/AMPAD_Lineage/blob/paper_rewrites_1/TCX_GenerateMonocleDS_new.R, and code used
354	to infer the lineage in ROSMAP is available here <u>https://github.com/Sage-</u>
355	Bionetworks/AMPAD_Lineage/blob/paper_rewrites_1/DLPFC_GenerateMonocleDS_new.R.
356	
357	Branch assignment and pseudotime calculation for samples
358	Branch assignment and pseudotime calculation was also performed using the Monocle package using
359	techniques described previously ¹⁹ . Briefly, pseudotime is calculated by first identifying a root point on
360	one of the two ends of the maximum diameter path in the tree. Then the pseudotime of each point is
361	calculated by projecting it to its closest point on the spanning tree and calculating the geodesic distance to
362	the root point. Assigning samples to branches is done by first identifying the branches of the spanning
363	tree and then assigning samples to the branch on which their projection to the spanning tree lies on.
364	
365	Association of pseudotime with AD status, hallmarks of Alzheimer's disease, and cognitive diagnosis
366	We test for association between disease pseudotime and AD case or control status with logistic regression
367	with AD case or control status as the outcome and inferred pseudotime as the dependent variable in both
368	the Mayo and ROS/MAP studies. We test for association between pseudotime and hallmarks of disease
369	in the ROS/MAP studies for both Braak (measure of tau pathology) score and CERAD score (measure of
370	amyloid pathology) with an ordinal logistic regression model, with the neuropath score as the ordered
371	outcome, and pseudotime as the dependent variable. Finally, we test for association between disease
372	pseudotime and cognitive diagnosis for the following ordered clinical diagnoses of no cognitive
373	impairment, mild cognitive impairment, and probable Alzheimer's disease with an ordinal logistic

374 regression model. All code for running these association tests is available <u>https://github.com/Sage-</u>

375 <u>Bionetworks/AMPAD_Lineage/blob/paper_rewrites_1/paper_figures.Rmd.</u>

376

377 Inferring cell type specific expression patterns given marker gene expression as a function of pseudotime 378 List of marker genes for different major cell types in the brain was curated from a previously published brain cell expression signature study³². The marker gene list was then pruned to include only genes that 379 were included in lineage construction. Each gene's expression as a function of pseudotime was then 380 381 obtained by smoothing using a smoothing spline of degree of freedom = 3 and normalized to lie in [0,1]. 382 The smoothing was done to remove the effects of technical noise introduced due to RNA-Seq and the 383 normalization was done since the absolute expression levels of genes might be very different from each 384 other. The smoothed and normalized expression of marker genes for each category were then averaged to 385 obtain the average marker gene expression as a function of pseudotime. A linear model was used to test 386 for association between average expression of a given cell type expression signature and pseudotime. 387

388 Association between GWAS loci and correlation with pseudotime

389 To test for association between pseudotime and LOAD GWAS genes, we computed the Spearman's 390 correlation between each gene's expression and pseudotime in the Mayo and ROS/MAP studies. Next, 391 we identified a set of genes implicated in AD GWAS loci in the International Genetics of Alzheimer's Project (IGAP)²³. We treated the set of genes described in Tables 1-3 of that study as high quality 392 393 candidate AD GWAS genes²³. We test for a difference between the correlation with pseudotime of 394 background of all other genes and the IGAP AD genes using a linear model, and see a significant increase 395 in correlation between gene expression and pseudotime in both the Mayo and ROS/MAP study for AD 396 GWAS genes.

397

398 Branch specific differential expression analysis

- 399 We perform a state specific differential expression analysis using a one-way ANOVA model in both the
- 400 Mayo and ROS/MAP studies. The branch with the highest proportion of AD controls is defined as the
- 401 reference branch for all analyses. We use Tukey's honest significant difference method to compute P-
- 402 values for the test for change in expression of a given gene compared to the reference branch. Genes are
- 403 grouped based on their branch and direction of change in expression for further downstream pathway
- 404 enrichment analyses. Code to run analyses are available here <u>https://github.com/Sage-</u>
- 405 <u>Bionetworks/AMPAD_Lineage/blob/paper_rewrites_1/DLPFC_DE_Anova.R</u> for ROS/MAP and here
- 406 <u>https://github.com/Sage-Bionetworks/AMPAD_Lineage/blob/paper_rewrites_1/TCX_DE_Anova.R</u> for
- 407 Mayo.
- 408
- 409 *Estimating branch specific gene expression signatures*
- 410 Branch specific expression signature was obtained by first calculating the average normalized expression
- 411 for all genes in each state/branch. This was followed by performing a bi-clustering using the pheatmap
- 412 package in R which uses hierarchical clustering on both samples and genes. We also used the pheatmap
- 413 package to visualize the state specific expression signatures.
- 414
- 415 *Gene set enrichment analyses*
- 416 For each branch specific differential expression gene set (DEGs) in both Mayo and ROS/MAP we
- 417 perform a gene set enrichment analysis against Gene Ontology pathways using the enrichR R package.
- 418 Only pathways with FDR < 0.05 are reported. The code we used to run the ROS/MAP DEG enrichments
- 419 are available here <u>https://github.com/Sage-</u>
- 420 <u>Bionetworks/AMPAD_Lineage/blob/paper_rewrites_1/lineage.Rmd</u>, the code we used to run the Mayo
- 421 DEG enrichments are available here <u>https://github.com/Sage-</u>
- 422 <u>Bionetworks/AMPAD_Lineage/blob/paper_rewrites_1/lineageTCX.Rmd</u>, and the code we used to run the
- 423 branch specific gene expression signature pathway enrichments is available here <u>https://github.com/Sage-</u>
- 424 <u>Bionetworks/AMPAD_Lineage/blob/paper_rewrites_1/resilience.Rmd</u>.

425

426 Whole-genome sequencing

427 Whole-genome sequencing was performed at the New York Genome Center for all individuals from the 428 ROS/MAP and Mayo cohorts. Detailed information for both data sets can be accessed via synapse 429 (DOI:10.7303/syn2580853). Briefly, 650ng of genomic DNA from whole blood was sheared using a 430 Covaris LE220 sonicator. DNA fragments underwent bead-based size selection and were subsequently 431 end-repaired, adenylated, and ligated to Illumina sequencing adapters. Libraries were sequenced on an 432 Illumina HiSeq X sequencer using 2 x 150bp cycles. Paired-end reads were aligned to the GRCh37 433 (hg19) human reference genome using the Burrows-Wheeler Aligner (BWA-MEM v0.7.8) and processed using the GATK best-practices workflow ^{53,54}. This included marking of duplicate reads by the use of 434 435 Picard tools v1.83, local realignment around indels, and base quality score recalibration (BOSR) via 436 Genome Analysis Toolkit (GATK v3.4.0). Joint variant calling files (vcfs) for whole-genome sequencing 437 data for the Mayo and ROS/MAP cohort were obtained through the AMP-AD knowledge portal 438 (www.synapse.org/#!Synapse:syn10901595). 439

440 Single variant association with pseudotime in two independent cohorts

441 Likelihood ratio tests within a linear regression framework were used to model the relationship between 442 the quantitative expression trait pseudotime and genetic variants in 436 AD cases. Genome-wide genetic 443 association analysis was performed for 305 female patients in the ROS/MAP cohort and 131 female 444 patients in the Mayo cohort for which both genotyping and post-mortem RNA-seq data was available. An 445 efficient mixed model approach, implemented in the EMMAX software suite, was used to account for potential biases and cryptic relatedness among individuals 55 . Only variants with MAF > 0.05, genotyping 446 call rates > 95%, minimum sequencing depth of 20 reads and Hardy-Weinberg equilibrium $p > 10^{-4}$ were 447 448 considered for analysis. Quantile-quantile plots (Figure S8-S9) for the test statistics showed no 449 significant derivation between expected and observed p-values, highlighting that there is no consistent 450 differences across cases and controls except for the small number of significantly associated variants.

- 451 Furthermore, the genomic inflation factor (lambda) was determined to be 0.99 for the Mayo and 0.98 for
- 452 the ROS/MAP single variant association tests. This highlights that potential confounding factors, such as
- 453 population stratification have been adequately controlled.

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597 Author Contributions

- 598 S.M. and B.A.L. designed the study. S.M. and B.A.L. performed the analyses. S.M., C.P., S.J., G.G.,
- 599 A.K.G., S.K.S., P.L.D.J., N.E.T., G.W.C., L.M.M., and B.A.L. wrote the manuscript.

601 Tables

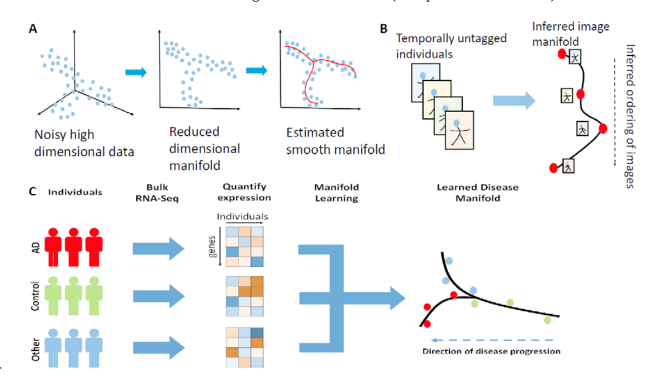
602 **Table 1** - Representative significant Gene Ontology pathway enrichments (FDR<0.05) of differentially

- 603 expressed genes for each branch (FDR < 0.05). Differential expressed genes are identified with an
- ANOVA analysis, with Branch 1 as the reference.

Brain Region	Direction	Branch	Representative Enriched Gene Ontology Terms
тсх	Down	2	prespliceosome (GO:0071010), mitochondrial electron transport, cytochrome c to oxygen (GO: 0006123)
		3	negative regulation of microtubule polymerization or depolymerization (GO:0031111)
		4	mitochondrial electron transport, NADH to ubiquinone (GO: 0006120), spliceosomal tri-snRNP complex (GO:0097526), negative regulation of microtubule depolymerization (GO:0007026)
		5	axon (GO:0030424), protein kinase C activity (GO:0004697),
		6	gamma-tubulin large complex (GO:0000931), U1 snRNP (GO:0005685), mitochondrial respiratory chain complex IV (GO:0005751), response to cadmium ion (GO:0046686)
	Up	2	
		3	fatty acid elongase activity (GO:0009922), ubiquitin protein ligase activity (GO:0061630)
		4	transforming growth factor beta-activated receptor activity (GO:0005024), hippo signaling (GO:0035329), regulation of extrinsic apoptotic signaling pathway via death domain receptors (GO: 1902041), regulation of DNA repair (GO: 0006282)
		5	regulation of apoptotic process (GO:0042981), leptin mediated signaling pathway (GO:0033210), negative regulation of hippo signaling (GO:0035331), small GTPase binding (GO:0031267)
		6	extracellular ligand-gated ion channel activity (GO:0005230), integral component of mitochondrial inner membrane (GO:0031305)
DLPFC	Down	2	DNA repair (GO:0006281), intracellular protein transport (GO:0006886)
		3	mismatch repair complex binding (GO:0032404)
		4	
		5	mitochondrial respiratory chain complex assembly (GO: 0033108)
		6	
	Up	2	racemase and epimerase activity (GO: 0016857)
		3	racemase and epimerase activity (GO: 0016857)
		4	vesicle mediated transport (GO: 0016192)
		5	NuRD complex (GO: 0016581)
		6	microtuble motor activity (GO:0003777), AP-2 adaptor complex binding (GO:0035612)

605 Figures

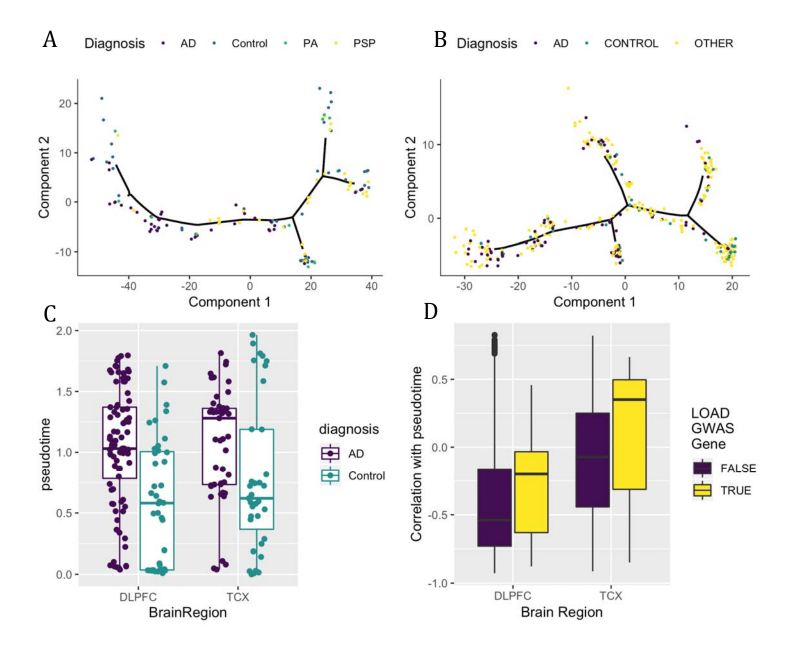
Figure 1 - Overview of manifold learning for unraveling staging in Alzheimer's disease. A) Illustration of
steps in manifold learning. B) A common application of manifold learning used in computer vision to
order temporally untagged images into sequences. C) Illustration of lineage inference process for LOAD.
RNA-seq samples with different disease diagnoses were pooled, batch normalized, and a smooth
manifold was learned for each brain region across individuals (each point is an individual).



612 Figure 2 - Manifold learning accurately infers disease states and stages from RNA-seq samples. A)

Estimated disease progression trees from temporal cortex (TCX) and B) dorsolateral prefrontal cortex

- 614 (DLPFC) brain regions showing localization of identified LOAD samples on particular branches. C)
- 615 Distribution of pseudotime for AD cases and controls for both DLPFC and TCX. D) Distribution of
- 616 expression correlation with pseudotime for both LOAD GWAS genes and non-LOAD GWAS genes.



- **Figure 3** Manifold learning replicates existing measures of staging in LOAD in DLPFC samples. A)
- 618 Samples colored by 3 different external measures of LOAD staging namely: Braak Score (tau pathology),
- 619 CERAD Score (amyloid pathology) and Cognitive Diagnosis (Clinical measure of disease severity).
- 620 Black lines denote inferred lineages. B) Distribution of samples by inferred stage for different distinct
- 621 stages in each of the three methods of measuring LOAD severity. Inferred disease stages generally
- 622 corresponded with all methods, and Cognitive diagnosis demonstrated the strongest alignment.

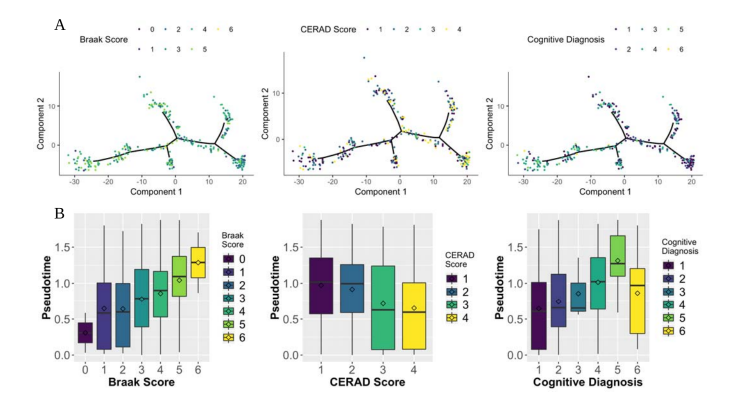


Figure 4 – Disease resistant state. A) The inferred manifold from the TCX region with samples colored
by their inferred disease subtype/state. State 5 (dots, circled) lies at the late end of the disease trajectory,
indicating a strong disease-like transcriptomic phenotype, yet most samples in the group did not have
pathologically diagnosed AD (Figure 2A). We hypothesize this group represents a disease resistant state
to the disease. B) Biclustering results of average expression from each disease state, with increased
expression of a gene cluster unique to State 5.

