1 Title: Molecular and cellular similarities in the brain of SARS-CoV-2 and Alzheimer's

2 disease individuals

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26 Abstract

27	Infection with the etiological agent of COVID-19, SARS-CoV-2, appears capable of
28	impacting cognition, which some patients with Post-acute Sequelae of SARS-CoV-2
29	(PASC). To evaluate neuro-pathophysiological consequences of SARS-CoV-2 infection,
30	we examine transcriptional and cellular signatures in the Broadman area 9 (BA9) of the
31	frontal cortex and the hippocampal formation (HF) in SARS-CoV-2, Alzheimer's disease
32	(AD) and SARS-CoV-2 infected AD individuals, compared to age- and gender-matched
33	neurological cases. Here we show similar alterations of neuroinflammation and blood-
34	brain barrier integrity in SARS-CoV-2, AD, and SARS-CoV-2 infected AD individuals.
35	Distribution of microglial changes reflected by the increase of Iba-1 reveal nodular
36	morphological alterations in SARS-CoV-2 infected AD individuals. Similarly, HIF-1 α is
37	significantly upregulated in the context of SARS-CoV-2 infection in the same brain
38	regions regardless of AD status. The finding may help to inform decision-making
39	regarding therapeutic treatments in patients with neuro-PASC, especially those at
40	increased risk of developing AD.
41	
42	Teaser
13	SARS-CoV-2 and Alzheimer's disease share similar neuroinflammatory processes which

43 SARS-CoV-2 and Alzheimer's disease share similar neuroinflammatory processes, which
44 may help explain neuro-PASC.

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- 46

48 Introduction

The consequences of SARS-CoV-2 infection have been well-studied in the respiratory 49 system; however, much less information is available regarding the neurological 50 consequences of infection (1). Previous evidence tentatively suggests that SARS-CoV-2 51 may be neuroinvasive, leading to a vast array of neurological symptoms, including 52 anosmia, ageusia, cognitive functions, and cerebrovascular disorders (2, 3). Neurological 53 complications of SARS-CoV-2 infection manifest with increased severity related to age 54 and shared medical history of metabolic disorders and other vascular risk factors (4–6). 55 Some strains within the coronavirus family are implicated in neuronal degeneration such 56 as HCoV-OC43, which can lead to glutamate excitotoxicity and neuronal degradation in 57 mice through cytokine production (7, 8). Because SARS-CoV-2 is a new addition to the 58 coronavirus family, little information exists regarding potential long-term neurological 59 complications that may impact COVID-19 survivors. This may be especially concerning 60 for aging individuals with increased vulnerability to developing age-related 61 neurodegenerative disease. 62

63

While it is currently unknown if neurological manifestations of SARS-CoV-2 infection 64 arise solely from systemic inflammation in the periphery or brain infiltration, it is well 65 established that secretion of cytokines and chemokines from the periphery allows the 66 recruitment of leukocytes and other cells to specific tissues (9, 10). This type of immune 67 response is speculated to accelerate blood-brain barrier (BBB) disruption and may 68 potentially damage cells within the central nervous system (CNS) (11). Such effects are 69 similar to neuropathology seen in Alzheimer's disease (AD) and other neurodegenerative 70 71 disorders, where leukocyte infiltration, BBB dysregulation, and microglial activation are observed. Thus, it is possible that SARS-CoV-2 may accelerate the onset and severity of 72

73	cognitive decline or AD in susceptible individuals (12, 13). Understanding the impact of
74	SARS-CoV-2 infection on the CNS and subsequent mechanisms associated with cognition
75	is particularly pressing, given the number of individuals presenting with neurological
76	symptoms of post-acute sequelae of SARS-CoV-2 (neuro-PASC), colloquially termed
77	'long-COVID' (14, 15).
78	
79	Here, we investigate potential brain mechanisms associated with SARS-CoV-2 infection
80	in SARS-CoV-2, AD and SARS-CoV-2 infected AD individuals by comparing
81	transcriptional and cellular responses in the cortical Broadman area 9 (cortical BA9) of the
82	frontal cortex and the hippocampal formation (HF); two brain regions deeply involved in
83	cognitive and emotional functions, to age- and gender-matched neurological cases. Here
84	we report evidence suggesting that SARS-CoV-2 infection promotes similar
85	pathophysiological features found in AD cortical BA9 and HF regions and possibly
86	exacerbate pre-existing AD pathophysiology.
87	
88	Results
89	Demographics of human postmortem cases
90	We examined postmortem tissue samples of cortical Broadman area 9 (cortical BA9) of
91	the frontal cortex and the hippocampal formation (HF) collected by the Neuropathology
92	Brain Bank and Research CoRE at Mount Sinai from 4 different groups: SARS-CoV-2,
93	AD, and AD individuals which became infected by SARS-CoV-2, compared to age- and
94	gender-matched neurological cases (Table S1). Each group comprises an equal number of
95	age-matched male and female patients, ensuring equal distribution with an average age of
96	79.6 among groups. Medical records indicate that all AD individuals have postmortem
97	ABC scores indicative of AD (Table S2), where A is a measure of amyloid beta

98	deposition, B is a measure of neurofibrillary degeneration based on the Braak and Brook
99	score, and C is scored based on neuritic plaques outlined by the Consortium to Establish a
100	Registry for Alzheimer's Disease diagnosis (CERAD) (16). SARS-CoV-2 infection was
101	confirmed by diagnostic polymerase chain reaction (PCR). Within the SARS-CoV-2 and
102	SARS-CoV-2 infected AD groups, all patients were symptomatic, admitted to the hospital,
103	and received oxygen supplementation by ventilator or cannula with disease onset to death
104	occurring on average, 27 days after diagnosis for patients with SARS-CoV-2 only and 32
105	days after diagnosis in infected AD patients (Table S1 and S2). Blood specimens were
106	collected and indicate several laboratory features of severe COVID-19 (Table S2), such as
107	increased C-reactive protein (CRP) and interleukin-6 (IL-6) (17).
108	
109	Gene expression of SARS-CoV-2 and AD groups compared to control cases in
110	cortical BA9 and HF tissues
111	The transcriptional profiles of AD and SARS-COV-2 infected postmortem cases from the
112	cortical BA9 were assessed by pairwise comparison to determine gene expression
113	compared to neurological control cases (Fig. 1A-C). Volcano plot distributions of gene
114	transcripts of SARS-CoV-2 and AD cases compared to neurological controls (Fig. 1A, B)
115	both reveal 39,901 genes which were used to determine differentially expressed genes
116	based on a nominal p-value less than 0.01 and an absolute log fold-change (logFC) greater
117	than 1. We find in the AD group, 287 genes are upregulated, and 131 genes are
118	downregulated. In the SARS-CoV-2 group, 179 genes are upregulated, and 426 genes are
119	downregulated. The Volcano plot distribution of gene transcripts of SARS-CoV-2 infected
120	AD cases compared to neurological controls (Fig. 1C) revealed from 38,021 genes, and
121	show 813 genes upregulated, 611 genes downregulated using the same cutoff for
122	significance as in (Fig. 1A, B).

124	Transcriptional profiles of AD and SARS-COV-2 cases from the HF were assessed
125	(p<0.01 and logFC>1) to determine gene expression, as compared to neurological control
126	cases (Fig. 1D-F), using 39,901 genes for SARS-CoV-2 and AD cases (Fig. 1D, E), and
127	35,527 genes for the SARS-CoV-2 infected AD cases (Fig. 1F). We find in the AD group,
128	304 genes upregulated, 214 genes downregulated. In the SARS-CoV-2 group, 230 genes
129	are upregulated, and 464 genes are downregulated. Additionally, in the SARS-CoV-2
130	infected AD group 466 genes upregulated, 582 genes downregulated.
131	
132	Similarity of gene expression in cortical BA9 and HF regions
133	In order to determine the relative similarity of gene expression changes induced by disease
134	state, a collection of genes known to be differentially expressed due to either SARS-CoV-
135	2 infection or AD were examined across three individual differential comparisons (Fig.
136	2A, 2B). While all three comparisons (AD versus Control, SARS-CoV-2 versus Control,
137	and SARS-CoV-2 infected AD versus Control) showed generally the same trends, the AD
138	and SARS-CoV-2 infected AD comparisons were strikingly similar, in both BA9 (Fig.
139	2A) and HF (Fig. 2B).
140	
141	Rank-rank hypergeometric overlap (RRHO) analysis comparing SARS-CoV-2 and AD
142	cases also reveals a positive correlation between the AD and SARS-CoV-2 groups in the
143	cortical BA9 (Fig. S1) as well as in the HF (Fig. S2), although to a lesser extent than in the
144	cortical BA9. We then filtered genes using Ingenuity Pathway Analysis software (IPA) to
145	determine the number of shared differentially regulated genes among the SARS-CoV-2,
146	AD, and SARS-CoV-2 infected AD groups within each tissue region (p<0.05 and absolute
147	z > 0.0001). Comparing these shared genes within the cortical BA9, we find that SARS-

148	CoV-2, AD and SARS-CoV-2 infected AD individuals shared 410 upregulated genes, 269
149	downregulated genes from 679 expressed genes (Fig. 2C; Data S1). In the HF we found
150	the three groups share 211 upregulated genes, 148 downregulated genes from 359
151	differentially expressed genes (Fig. 2D; Data S2).
152	
153	Pathway activation of SARS-CoV-2 and AD groups
154	IPA was employed to further characterize pathways in SARS-CoV-2, AD and SARS-
155	CoV-2 infected AD groups compared to the neurological controls. Within the cortical
156	BA9 region (Fig. 3A), for example several pathways such as neuroinflammation, TREM1,
157	and cell senescence show increased activation in the SARS-CoV-2, AD and SARS-CoV-2
158	infected AD groups. TREM1 signaling, the most highly activated pathway in this dataset
159	(Fig. 3A, B), is expressed primarily on myeloid cells, such as macrophages and microglia,
160	and is involved in pro-inflammatory immune responses (18). Although cellular
161	senescence due to age is a natural process where telomeres shorten over time, this process
162	also occurs during cellular stress due to inflammation, including from viral infections,
163	leading to a senescence-associated secretory phenotype (SASP) such as metalloproteinases
164	(MMP's), and inflammatory cytokines (19, 20) (Fig. 3A). Another pathway of interest,
165	SNARE, shows decreased activation in AD and SARS-CoV-2 groups (Fig. 3A), with the
166	most significant reduction seen in the SARS-CoV-2 infected AD cases group. SNARE
167	proteins play an essential role in neurotransmitter release, and altered function is
168	implicated in the pathophysiology of neurodegenerative diseases such as AD, where
169	SNARE proteins affect β -amyloid (A β) accumulation and cytoplasmic transport of
170	neurofibrillary tangles (21).

172	Within the HF (Fig. 4A), for example interleukin-8 (IL-8) another neuroinflammatory
173	pathway, and ciliary neurotrophic factor (CNTF) signaling is upregulated in SARS-CoV-
174	2, AD and SARS-CoV-2 infected AD groups, while cAMP response element-binding
175	protein (CREB) is also upregulated in SARS-CoV-2 groups but downregulated in AD-
176	only (Fig. 4A). IL-8 has several important roles, including endothelial cell migration and
177	chemoattraction of neutrophils (22). One way IL-8 aids cell migration is by enhancing the
178	expression of molecules such as MMP-2, MMP-9, involved in BBB integrity and
179	induction of neuronal apoptosis, and VEGF-A, involved in vascular permeability and
180	angiogenesis, thus having a potential effect on vascular damage (23–25). CNTF signaling
181	aids in the prevention of neuronal degeneration after injury and is neuroprotective in
182	diseases such as multiple sclerosis (MS) and amyotrophic lateral sclerosis (ALS) (26).
183	Interestingly, CREB signaling modulates processes in consolidating memory and
184	information processing and is inhibited in AD (27). Consistent with these findings, we
185	also see a reduction in CREB signaling in the AD individuals; however, this effect is
186	reversed in SARS-CoV-2 infected AD individuals in the HF. It is important to note that
187	TREM1 signaling is upregulated in two of the three sample groups (SARS-CoV-2 and
188	SARS-CoV-2 infected AD cases), but no predictions for this pathway occurs in the AD
189	group. As such, this pathway is not included in the analysis within the HF (Fig 4A,B).
100	

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Additionally, upregulation of integrin signaling and nerve growth factor (NGF) signaling occur in both cortical BA9 and HF tissue (**Fig's. 3B and B4**) in all SARS-CoV-2, AD and SARS-CoV-2 infected AD groups. Neuronal function shown by the upregulation of NGF signaling is critical for the survival of neurons, and alterations of NGF signaling has been implicated in neurodegenerative disorders such as AD (28). Integrin signaling is used for a

196	diverse array of functions within the CNS via cell-to-cell and cell-to-extracellular matrix
197	interactions (29).

egulation of inflammatory microglia responses in cortical BA9 and HF sections
egion-matched formalin-fixed paraffin-embedded (FFPE) tissue from the contralateral
emisphere of cortical BA9 and HF used for transcriptional studies underwent
nmunohistochemical (IHC) analysis for evidence of microglial activation (Iba-1), the
resence of SARS-CoV-2 (SARS-CoV-2 nucleocapsid), and vascular integrity, assessed
a hypoxia-inducible factor-1α subunit (HIF-1α).
euroinflammation was assessed with a pan-microglia marker, Iba-1, which is
pregulated by microglia in the context of inflammation and reveals morphological
terations associated with their activation state (Fig. 5). Compared to neurological
ontrols, microglia in AD, SARS-CoV-2, and SARS-CoV-2 infected AD cases are more
amerous and display an activated phenotype, with retracted, thickened processes and
nlarged somas (Fig. 5 A-D). Microglia also appear to accumulate around blood vessels in
e context of AD and SARS-CoV-2 groups but not in unaffected neurological controls
rain (Fig. 5 E-H), suggesting factors at the level of the BBB may participate in
icroglial activation. Microglial nodules, which are commonly observed in
euroinflammatory disease, were seen in all conditions but appeared larger and more
equently in infection, as compared to AD-only and controls (Fig. 5 I-L). When
etectable, SARS-CoV-2 appears to be restricted to the endothelium (Fig. 5 G, H, K, L)
nd did not colocalize with Iba-1 or glial fibrillary acidic protein (GFAP; data not shown),
aggesting neither microglia nor astrocytes harbor productive virus. Nonbiased
antitation revealed an overall increase in the number of microglia in the HF and cortical

221	BA9 of patients with AD, SARS-CoV-2, and SARS-CoV-2 infected AD cases. A
222	statistically significant increase in the number of microglia occurs in AD alone in the HF
223	and SARS-CoV-2 infected AD cases in both the HF and cortical BA9 (Fig. 5). An
224	increase in the frequency of nodular lesions was seen with SARS-CoV-2 infection, with
225	and without AD, in both brain regions assessed. However, a statistically significant
226	increase in nodular lesions are present in SARS-CoV-2 infected AD cases (Fig. 5P-R).
227	
228	Regulation of vascular damage using hypoxia inducible factor- 1α (HIF-1 α) detection
229	Tissues were further assessed for possible hypoxia by IHC using an antibody against the $\boldsymbol{\alpha}$
230	subunit of HIF-1, which is stabilized under hypoxic conditions. Upregulation and
231	stabilization of HIF-1 α is most pronounced in the context of SARS-CoV-2 infection,
232	regardless of AD status in cortical BA9 (Fig. 6A-D) and HF (Fig. 6E-H). Nonbiased area
233	positivity quantitation revealed HIF-1 α is only slightly elevated in cortical BA9 of AD
234	patients, as compared to age-matched neurological controls (Fig. 6J). In contrast, a greater
235	increase occurs in the HF from SARS-CoV-2 individuals, with and without AD. SARS-
236	CoV-2 only samples showed a significant increase in HIF-1 α of cortical BA9 when
237	compared to controls (Fig. 6I-K).
238	
239	Discussion
240	The finding from our study suggests that SARS-CoV-2 and AD infected individuals
241	share similar alterations of regulatory patterns of immune-inflammatory pathways and
242	pathways involved with cognition as suggested by recent meta-analyses showing shared
243	neuroinflammation and microvascular injury, in particular AD and SARS-CoV-2
244	infected individuals (30). Additionally, Zhou and colleagues found a significant overlap
245	in cerebrospinal fluid (CSF) monocytes and markers in AD and COVID-19, which also

246	occurs in our dataset (30). The similarities of transcriptional profiles and cellular
247	pathophysiology in SARS-CoV-2 and SARS-CoV-2 infected individuals support the
248	potential role of SARS-CoV-2 infection on the CNS, leading to neuro-PASC symptoms
249	such as brain fog and memory loss.
250	
251	Notably, our study identifies similar neuroinflammatory profiles in SARS-CoV-2, AD and
252	AD SARS-CoV-2 infected individuals at the transcriptional and cellular levels in both the
253	cortical BA9 and HF brain regions. This suggests that SARS-CoV-2 generates a similar
254	neuroinflammatory environment in neurodegenerative disorders like AD. This was
255	highlighted by the regulation of TREM1, neuroinflammation, and cellular
256	senescence/inflammatory pathways present in all groups and further established by the
257	widespread microglial activation seen in AD and SARS-CoV-2 infected AD cases, and in
258	particular, the nodular formation seen in SARS-CoV-2 infected AD cases. Microglia
259	nodule formation is present in some neurodegenerative diseases, such as MS, and viral
260	infections, such as herpes simplex virus (HSV) and human immunodeficiency virus (HIV)
261	(31, 32). A potentially compounding finding is the nodular formation that is initially
262	characterized by abundant presence of activated microglia and innate immune factors
263	leading to toll-like receptor (TLR) signaling and upregulations of inflammasome genes,
264	leading to T cell stimulation and ultimately the destruction of neurons (31) . This may be
265	of particular concern in the SARS-CoV-2 infected AD cases, where we primarily observe
266	increased nodular lesions. This suggests SARS-CoV-2 further promotes
267	neuroinflammation in AD, which likely advances the progression and severity of CNS
268	disease in these individuals. Interestingly, a large retrospective study of patients 65 years
269	or older revealed that patients with SARS-CoV-2 were at an increased risk for a new AD
270	diagnosis within a year of their SARS-CoV-2 diagnosis, with the most significant risk

271	seen among those 85 years and older (33) . This may underscore the critical role of
272	preexisting inflammation in the brain, which is seen in the context of 'normal' aging, in
273	promoting or advancing AD progression.

274

Our findings also support the notion that SARS-CoV-2 may cause cognitive deficits via 275 regulation of pathways associated with cognition and neuroinflammation. Here, we show 276 changes in the transcriptional regulation of SNARE and NGF pathways, suggesting 277 impaired neuronal health and function, presumably negatively impacting cognitive 278 function. We also observed potential damage to the vasculature via increased regulation of 279 HIF-1 α , integrin signaling, and IL-8 signaling. It is important to note a possibility of 280 vascular damage due to ventilation prior to death in SARS-CoV-2 infected individuals; 281 however, vascular damage by SARS-CoV-2, as assessed by HIF-1a, is observed in non-282 human primates that were euthanized at designated end-of-life time points precluding 283 breathing intervention (34). These findings may point to a possible route for lymphocyte 284 transmigration following chemoattractant gradients such as IL-8 and enhancement of 285 MMPs and VEGF, which are aided by integrins such as cellular adhesion molecules such 286 as ICAM-1 and are implicated with inflammatory signal transduction (35, 36). This 287 process leads to T cell activation, which may be responsible for the observed 288 inflammatory environment. Our transcriptional data showed an abundant upregulation of 289 TLRs (TLRs 1, 5, 7, and 8) related to the inflammasome. Further analysis of this dataset 290 also indicated the potential role for T cells through IL-7 pathway activation and CD86 291 upregulation; however, we did not confirm changes in the number of T cells, and no 292 evidence of leukocyte infiltration into the CNS compartment was observed in any disease 293 state in this investigation. Future studies will be required to determine the role of 294 295 leukocytes in the observed pathophysiology.

296

297	Although neuroinflammation and vascular injury were prominent features of SARS-CoV-
298	2 and SARS-CoV-2 infected AD cases brain pathology, the direct role of the virus is
299	unclear. We did detect SARS-CoV-2 nucleocapsid in some SARS-CoV-2 brain tissues
300	that appear to be restricted to the vasculature. This finding is supported by other studies
301	suggesting SARS-CoV-2 is sporadically present in brain tissue (37). Importantly, we only
302	investigated CNS regions with the greatest significance in AD. These regions may be less
303	prone to SARS-CoV-2 infection than others, such as the olfactory bulb and tract, where
304	olfactory neurons are proposed to be infected through viral spread from olfactory
305	epithelium (38). Our findings of the viral nucleocapsid limited to the endothelium suggest
306	the hematological spread of SARS-CoV-2 to the CNS that may not extend to the neurons
307	in the cortical BA9 and HF yet is still capable of inducing widespread inflammation in the
308	brain.
309	
310	Even in the absence of a detectable virus in the neurons or neural cells, SARS-CoV-2
311	may impact cognitive dysfunction through TREM1 activation of the NLRP3
312	inflammasome and subsequent pyroptosis, where pro-caspase 1 cleavage and subsequent
313	cleavage and activation of IL-1 β , IL-18 and gasmerdin D pore formation in cells
314	ultimately leading to pyroptosis (39, 40). NLRP3 activation is reported in AD and
315	COVID-19, and is suggested by formation of microglial nodules demonstrated in these
316	cases (41–43).`
317	
318	SARS-CoV-2 is not the first virus to be implicated in cognitive dysfunction. This is a facet
319	shown with other viruses such as HIV, HSV, and Epstein-Barr virus (EBV) (44, 45).
220	While this study connect and dist the systems of disease any provision in COVID 10

320 While this study cannot predict the outcome of disease progression in COVID-19

321	survivors, the present findings that SARS-CoV-2 infection can recapitulate AD-type
322	transcriptional and cellular neuroinflammatory patterns among other in a very short time
323	frame makes it critical to understand how SARS-CoV-2 impacts long-term cognition. The
324	increase in nodular formation present in SARS-CoV-2 infected AD cases tissue also
325	demonstrates a critical need to functionally determine potentially synergistic effects of AD
326	and SARS-CoV-2. This is underscored by the prevalence of cognitive dysfunction seen
327	among neuro-PASC patients, making it imperative that the link between SARS-CoV-2
328	and cognition be intensively investigated to identify potential therapeutic strategies for
329	halting cognitive decline in these individuals. Collectively, our results demonstrate several
330	key areas of overlap between the neurological effects of SARS-CoV-2 infection and AD.
331	These findings may help inform decision-making regarding therapeutic treatments in
332	patients with COVID-19, especially those who may be at increased risk of developing AD.
333	
333 334	Materials and Methods
	Materials and Methods Patients
334	
334 335	Patients
334 335 336	Patients Brain tissue was collected from the cortical BA9 region (cortical BA9) and hippocampal
334335336337	Patients Brain tissue was collected from the cortical BA9 region (cortical BA9) and hippocampal formation (HF) of four SARS-CoV-2 cases, four Alzheimer's disease (AD) cases and four
 334 335 336 337 338 	Patients Brain tissue was collected from the cortical BA9 region (cortical BA9) and hippocampal formation (HF) of four SARS-CoV-2 cases, four Alzheimer's disease (AD) cases and four non-SARS-CoV-2 or AD autopsies. With each case, one hemisphere of the brain was
 334 335 336 337 338 339 	Patients Brain tissue was collected from the cortical BA9 region (cortical BA9) and hippocampal formation (HF) of four SARS-CoV-2 cases, four Alzheimer's disease (AD) cases and four non-SARS-CoV-2 or AD autopsies. With each case, one hemisphere of the brain was formalin-fixed paraffin-embedded (FFPE) and the other hemisphere was frozen to
 334 335 336 337 338 339 340 	Patients Brain tissue was collected from the cortical BA9 region (cortical BA9) and hippocampal formation (HF) of four SARS-CoV-2 cases, four Alzheimer's disease (AD) cases and four non-SARS-CoV-2 or AD autopsies. With each case, one hemisphere of the brain was formalin-fixed paraffin-embedded (FFPE) and the other hemisphere was frozen to generate fresh-frozen blocks. FFPE tissue was used for microscopy and region matched
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346 **RNA sequencing**

347	Samples collected and homogenized in RNAzol RT (Molecular Research Center, Inc.)
348	were then processed using the Zymo Clean and concentrator Kit (Zymo Research) to
349	collect total RNA following the manufacturers protocols. cDNA library construction and
350	sequencing were conducted by Genewiz (Azenta Life Sciences). Total RNA samples were
351	quantified using Qubit 2.0 Fluorometer (Life Technologies, Carlsbad, CA, USA) and
352	RNA integrity was checked with 4200 TapeStation (Agilent Technologies, Palo Alto, CA,
353	USA).
354	
355	Samples were treated with TURBO DNase (Thermo Fisher Scientific, Waltham, MA,
356	USA) to remove DNA contaminants, followed by rRNA depletion using QIAseq®
357	FastSelect TM –rRNA HMR kit (Qiagen, Germantown, MD, USA); conducted following
358	the manufacturer's protocol. RNA sequencing libraries were constructed with the
359	NEBNext Ultra II RNA Library Preparation Kit for Illumina by following the
360	manufacturer's recommendations, where enriched RNAs were fragmented for 15 minutes
361	at 94 °C. First strand and second strand cDNA are subsequently synthesized. cDNA
362	fragments are end repaired and adenylated at 3'ends, and universal adapters are ligated to
363	cDNA fragments, followed by index addition and library enrichment with limited cycle
364	PCR. Sequencing libraries were validated using the Agilent Tapestation 4200 (Agilent
365	Technologies, Palo Alto, CA, USA), and quantified using Qubit 2.0 Fluorometer
366	(ThermoFisher Scientific, Waltham, MA, USA) as well as by quantitative PCR (KAPA
367	Biosystems, Wilmington, MA, USA). The sequencing libraries were multiplexed and
368	clustered on one lane of a flowcell. After clustering, the flowcell was loaded on the
369	Illumina HiSeq 4000 instrument according to manufacturer's instructions. The samples
370	were sequenced using a 2x150 Pair-End (PE) configuration.

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372	Image analysis and base calling were conducted by the HiSeq Control Software (HCS).
373	Raw sequence data (.bcl files) generated from Illumina HiSeq was converted into FASTQ
374	files and de-multiplexed using Illumina's bcl2fastq 2.17 software. One mismatch was
375	allowed for index sequence identification. After investigating the quality of the raw data,
376	sequence reads were trimmed to remove possible adapter sequences and nucleotides with
377	poor quality using Trimmomatic v.0.36. The trimmed reads were mapped to the GRCh38
378	reference genome available on ENSEMBL using the STAR aligner v.2.5.2b. BAM files
379	were generated as a result of this step. Unique gene hit counts were calculated by using
380	feature Counts from the Subread package v.1.5.2. Only unique reads that fell within exon
381	regions were counted.
382	
383	Immunohistochemistry
383 384	Immunohistochemistry Tissue was cut at 6 µm on a Leica Microtome to for immunohistochemistry that was
384	Tissue was cut at 6 μ m on a Leica Microtome to for immunohistochemistry that was
384 385	Tissue was cut at $6 \mu m$ on a Leica Microtome to for immunohistochemistry that was performed formalin-fixed paraffin-embedded (FFPE) brain sections as previously
384 385 386	Tissue was cut at 6 µm on a Leica Microtome to for immunohistochemistry that was performed formalin-fixed paraffin-embedded (FFPE) brain sections as previously described (46) (Iba-1 staining was conducted on a Ventana Benchmark using OptiView
384 385 386 387	Tissue was cut at 6 µm on a Leica Microtome to for immunohistochemistry that was performed formalin-fixed paraffin-embedded (FFPE) brain sections as previously described (46) (Iba-1 staining was conducted on a Ventana Benchmark using OptiView and UltraView detection kits provided by Roche (Roche Molecular Systems, Inc).
384 385 386 387 388	Tissue was cut at 6 µm on a Leica Microtome to for immunohistochemistry that was performed formalin-fixed paraffin-embedded (FFPE) brain sections as previously described (46) (Iba-1 staining was conducted on a Ventana Benchmark using OptiView and UltraView detection kits provided by Roche (Roche Molecular Systems, Inc). Sections were deparaffinized in xylene and rehydrated through an ethanol series ending in
384 385 386 387 388 389	Tissue was cut at 6 µm on a Leica Microtome to for immunohistochemistry that was performed formalin-fixed paraffin-embedded (FFPE) brain sections as previously described (46) (Iba-1 staining was conducted on a Ventana Benchmark using OptiView and UltraView detection kits provided by Roche (Roche Molecular Systems, Inc). Sections were deparaffinized in xylene and rehydrated through an ethanol series ending in distilled water. Heat-mediated antigen retrieval was carried out in a vacuum oven with
384 385 386 387 388 389 390	Tissue was cut at 6 µm on a Leica Microtome to for immunohistochemistry that was performed formalin-fixed paraffin-embedded (FFPE) brain sections as previously described (46) (Iba-1 staining was conducted on a Ventana Benchmark using OptiView and UltraView detection kits provided by Roche (Roche Molecular Systems, Inc). Sections were deparaffinized in xylene and rehydrated through an ethanol series ending in distilled water. Heat-mediated antigen retrieval was carried out in a vacuum oven with Tris-EDTA buffer (10mM Trizma base, 1mM EDTA, 0.05% Tween 20, pH 9.0) or
 384 385 386 387 388 389 390 391 	Tissue was cut at 6 µm on a Leica Microtome to for immunohistochemistry that was performed formalin-fixed paraffin-embedded (FFPE) brain sections as previously described (46) (Iba-1 staining was conducted on a Ventana Benchmark using OptiView and UltraView detection kits provided by Roche (Roche Molecular Systems, Inc). Sections were deparaffinized in xylene and rehydrated through an ethanol series ending in distilled water. Heat-mediated antigen retrieval was carried out in a vacuum oven with Tris-EDTA buffer (10mM Trizma base, 1mM EDTA, 0.05% Tween 20, pH 9.0) or sodium citrate buffer (10mM sodium citrate, 0.05% Tween 20, pH 6.0). All washes were
384 385 386 387 388 389 390 391 392	Tissue was cut at 6 µm on a Leica Microtome to for immunohistochemistry that was performed formalin-fixed paraffin-embedded (FFPE) brain sections as previously described (46) (Iba-1 staining was conducted on a Ventana Benchmark using OptiView and UltraView detection kits provided by Roche (Roche Molecular Systems, Inc). Sections were deparaffinized in xylene and rehydrated through an ethanol series ending in distilled water. Heat-mediated antigen retrieval was carried out in a vacuum oven with Tris-EDTA buffer (10mM Trizma base, 1mM EDTA, 0.05% Tween 20, pH 9.0) or sodium citrate buffer (10mM sodium citrate, 0.05% Tween 20, pH 6.0). All washes were performed using tris buffered saline containing Tween 20 (TTBS; 0.1M Trizma base,

396	at room temperature and detected using the appropriate biotinylated secondary antibody
397	(1:200, Vector Labs, BA-1100, BA-2000) and alkaline phosphatase-Vector Red according
398	to manufacturer instructions (Vector Labs). Tissues were counterstained with Mayer's
399	hematoxylin and coverslipped.
400	
401	Imaging and quantitation
402	Slides were scanned with the Axio Scan.Z1 digital slide scanner (Zeiss). Brightfield
403	images were acquired using HALO (Indica Labs, v3.4.2986.151). Figures were created in
404	Photoshop (Adobe, v23.5.1) by brightness and contrast adjustments applied to the entire
405	image.
406	
407	Threshold and multiplex analyses were performed with HALO algorithms for non-biased
408	quantitation of proteins of interest without processing, as described previously (38). For
409	microglia quantitation, hematoxylin-stained nuclei were used to quantify the total number
410	of cells and those with Vector Red intensity above a rigorous threshold (Iba-1+).
411	Microglia frequency is reported as the percentage of total nuclei in the tissue section
412	assessed. To assess frequency of nodular lesions, all Iba-1-stained tissues were viewed in
413	HALO and nodular lesions made up of 3 or more Iba-1+ microglia in contact with one
414	another were counted as a single nodule. The total number of nodules for each tissue is
415	reported per tissue area. Quantitation of HIF-1 α was performed using an area
416	quantification algorithm for Vector Red intensity. Annotations were drawn to outline the
417	tissue and exclude empty spaces and glass. The annotated area was analyzed for overall
418	quantity of Vector Red positivity per micron ² of tissue. Two-tailed Mann-Whitney U tests
419	were performed with GraphPad Prism software, v9.3.1. Data are expressed as mean \pm
420	SEM. P values ≤ 0.05 are considered significant.

421

422 Bioinformatics Data Access and Analysis

423	Sample similarity was assessed with PCA analysis on VST-transformed expression values.
424	Genes were filtered to remove lowly expressed genes, defined as fewer than 5 samples
425	showing a minimum read count of 1 read, prior to performing differential analysis with
426	DESeq2, in R (4.2.0) (47). The transcriptional profiles of SARS-CoV-2 and AD
427	postmortem cases from the cortical BA9 and HF were assessed by pairwise comparison to
428	the neurologically healthy control cases. Genes were assigned as differentially expressed
429	if the nominal p-value was less than 0.05 and the absolute log2FC exceeded 1. An
430	additional round of differential expression testing was performed using a model containing
431	covariates (COVID status, Alzheimer's status, gender, and brain region), to extract the
432	transcriptional effects of the individual covariates. Biological pathways and key regulators
433	impacted by disease were identified using QIAGEN Ingenuity Pathway Analysis (IPA)
434	(QIAGEN Inc., Version 73620684) (48) Genes with a threshold of p<0.05 were used as
435	input for IPA. The relative similarities of transcriptional changes in the DESeq2
436	comparisons were assessed using Rank-rank hypergeometric overlap (RRHO2) analysis.
437	

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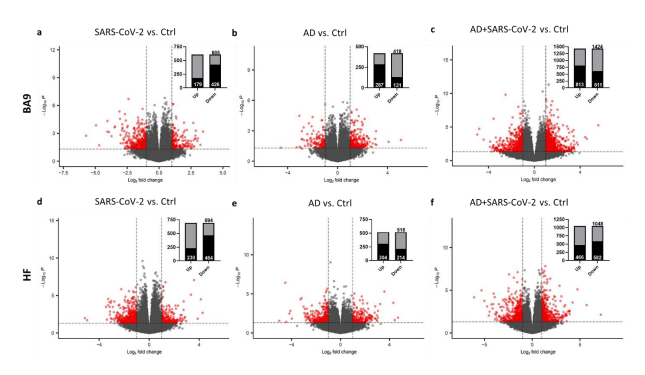
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614	Methodology: EG, TF, GP
615	Investigation: EG, KT, SN, LH, ME, GH, BM, NJ, TF
616	Visualization: EG, ME, LS, LH, TF
617	Supervision: TF, GMP
618	Writing—original draft: EG, ME, TF
619	Writing—review & editing: EG, SN, NJ, TF, GMP
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624	neuroimaging findings. The other authors declare that they have no conflict of interest
625	with the contents of this article.

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627	Data and materials availability: The authors declare that the data supporting the findings
628	of this study are available within the paper and its supplementary information files. The
629	raw data discussed in this publication will be accessible through NCBI's Gene Expression
630	Ominubus (GEO) upon publication (49).

631 Figures and Tables

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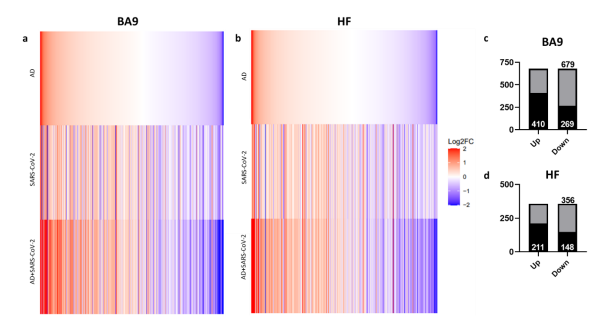
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Fig. 1. Gene expression in cortical BA9 and HF tissue.

Volcano plot distribution of gene transcripts of the cortical BA9 region in as shown in **A**, SARS-CoV-2 cases, in **B**, AD cases, and in **C**, SARS-CoV-2 infected AD cases compared to neurological control cases. The HF region in panels **D**, SARS-CoV-2 cases, **E**, AD cases, and **F** SARS-CoV-2 infected AD cases compared to neurological control cases. Volcano plots were generated from 39,901 genes in A, B, D and E, and 38,021 genes in C 35,527 genes in F. Transcripts with nominal p<0.01 and an absolute log fold-change (logFC) greater than 1 are indicated in red and shown in inlayed bar graphs wherein the number of upregulated and downregulated genes are indicated in black as part of the total of differentially regulated genes (grey).



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Fig. 2. Similarity of gene expression in cortical BA9 and HF tissue in SARS-CoV-2 and SARS-CoV-2 infected AD relative to AD individuals based on log2FC.

648 Genes known to be differentially expressed due to either SARS-CoV-2 infection or AD were

examined with respect to their gene expression changes, with the given gene set being selected for

cortical BA9, shown in panel **A**, or HF, shown in panel **B**. The log2FC of the genes for three

651 individual differential comparisons (AD versus Control, SARS-CoV-2 versus Control, and

652 SARS-CoV-2 infected AD versus Control) are shown for the BA9 and HF tissue as shown in 653 panels A and B, respectively. Then, using Ingenuity pathway analysis (IPA) as a filtering system

653 panels A and B, respectively. Then, using Ingenuity pathway analysis (IPA) as a filtering system 654 for genes within the database, we compared the similarity in expression of genes present in all

654 for genes within the database, we compared the similarity in expression of genes present in all 655 datasets (SARS-CoV-2, AD and SARS-CoV-2 infected AD cases), shown in panel **C** and **D**, in

656 both the cortical BA9 and HF.

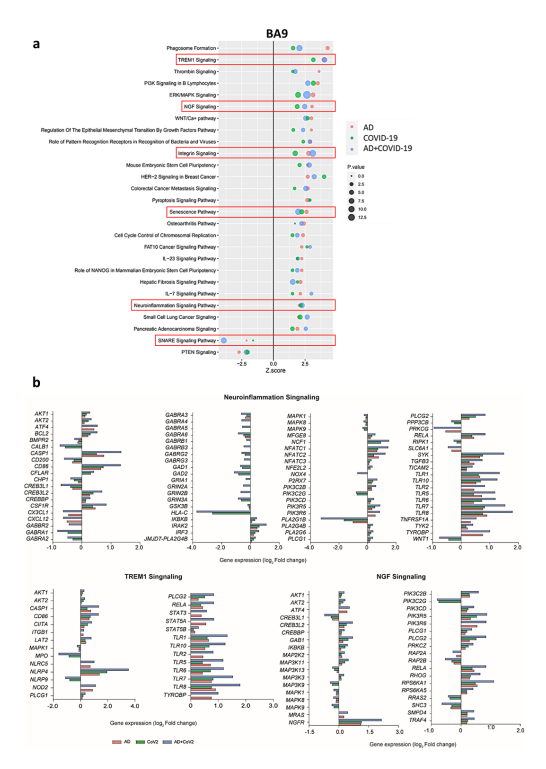
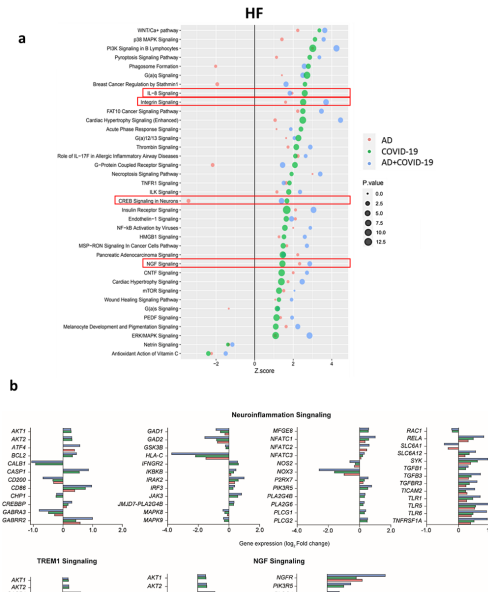


Fig. 3. Changes in Signaling Pathways within the cortical BA9.

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Canonical Pathway Similarities as seen in panel A, show top regulated canonical pathways 660 within the cortical BA9 in SARS-CoV-2, AD, and SARS-CoV-2 infected AD cases in 661 reference to the control. Activation score (Z-score) is shown on the X-axis, while the 662 pathways are indicated on the Y-axis. The color of the points indicates the IPA 663 comparison, while the size of the point represents the -log10 p-value of the IPA 664 comparison, with the larger points indicating the lowest p-values. The predicted gene 665 regulation of the Neuroinflammation, TREM1 and NGF pathways, shown in panel **B**, 666 indicate that SARS-CoV-2, AD and SARS-CoV-2 infected AD groups have similar 667 expression in key inflammatory and neuronal pathways. 668



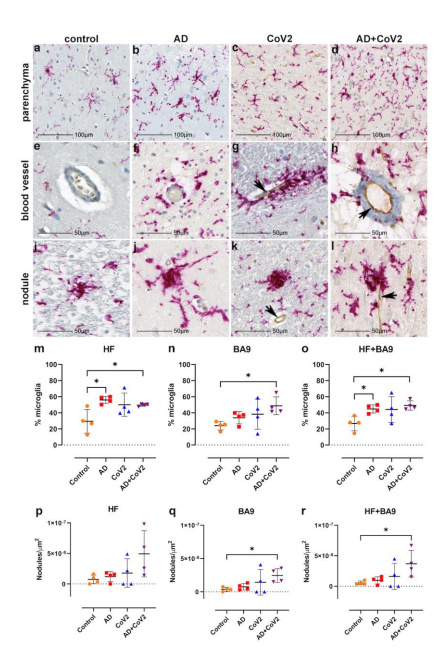
2.0

AKT1 AKT2 CASP1 CASP5 ATF4 PLCG1 CREBBE PLCG2 CD86 CIITA LAT2 RAC1 GAB1 IKRKR RAF1 RALA MAP2K2 NLRP9 PLCG1 MAP3K11 RELA PLCG2 MAP3K13 RPS6KA1 RELA RPS6KA5 MAPK8 TLR1 MAPK9 RRAS TLR5 MRAS TLR 2.0 -1.0 1.0 2.0 -1.0 1.0 3.0 1.5 -1.5 0.0 Gene expression (log, Fold change) Gene expression (log₂ Fold change) AD CoV2 AD+CoV2

Fig. 4. Changes in Signaling Pathways within the HF.

Canonical pathway similarities as seen in panel A, top regulated canonical pathways were 671 compared within the HF in SARS-CoV-2, AD, and SARS-CoV-2 infected AD cases in 672 reference to the control. Activation score (Z-score) is shown on the X-axis, while the 673 pathways are indicated on the Y-axis. The color of the points indicates the IPA 674 comparison, while the size of the point represents the -log10 p-value of the IPA 675 comparison, with the larger points indicating the lowest p-values. The predicted gene 676 regulation of the Neuroinflammation, TREM1 and NGF pathways, shown in panel **B**, 677 indicate that SARS-CoV-2, AD and SARS-CoV-2 infected AD groups have similar 678 expression in key inflammatory and neuronal pathways. 679

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Fig. 5. Microgliosis and nodular lesions in neurological controls, SARS-CoV-2, AD and SARS-CoV-2 infected individuals.

The level of microglial activation was assessed by immunohistochemical staining using anti-Iba-1 683 antibody with Vector Red. The presence of SARS-CoV-2 was assessed using an antibody against 684 the virus nucleocapsid and visualized with DAB. Parenchymal microglia are more frequent and 685 686 highly activated in the context of infection with and without AD, as shown by thickened processes, enlarged soma, and loss of individual cellular domain in panels **B-D**, as compared to 687 age-matched controls shown in A and E. When present, SARS-CoV-2 localizes to the blood 688 vessel endothelium shown in panels G, H, K, L (black arrows). Microglia appear to gather around 689 blood vessels in disease but do not form cuffs (F, G, H). Nodular lesions are seen in most cases 690 assessed, regardless of disease status (I-L), however, they appear larger and more frequent in the 691 context of disease shown in panels J-L, as compared to controls, panel I. A multiplex algorithm 692 was used to count all cells, using DAPI+ nuclei, with HALO and calculate percent frequency of 693 Iba-1+ microglia, shown in panels M-O. There was upregulation of Iba-1 in the hippocampus as 694 compared to the cortical BA9 region for most cases, though there is a significantly higher ratio of 695 microglia relative to other cells when comparing the AD and SARS-CoV-2 infected AD cases to 696

the control group shown in panel **M**. This trend seems to hold true for cortical BA9 region as

698 well, where the only groups with a significant difference in microglia ratios are the control and 699 SARS-CoV-2 infected AD cases shown in panel **N**. When all cases have both regions averaged,

the level of microgliosis is shown to be higher in the AD and SARS-CoV-2 infected AD cases

when compared to control in panel **O**. Graphs **P-R** show the normalized counts of microglial

nodule frequency. A higher frequency of nodules is seen in the HF, however, the difference

between groups did not reach statistical significance as shown in panel **P**. In contrast, fewer

nodules are seen in cortical BA9, overall. A statistically significant higher number of lesions are

seen in SARS-CoV-2 infected AD cases, as compared to control cortical BA9, suggesting greater
 inflammation in the cortical BA9 of patients with both AD and SARS-CoV-2 infection, shown in

707 panel **O**. Significance was maintained between the control and SARS-CoV-2 infected AD groups

when the average was taken for both brain regions per case shown in panel \mathbf{R} . Statistics were

performed with a two-tailed Mann-Whitney U test. *p<0.05. Data are expressed as mean \pm SEM.

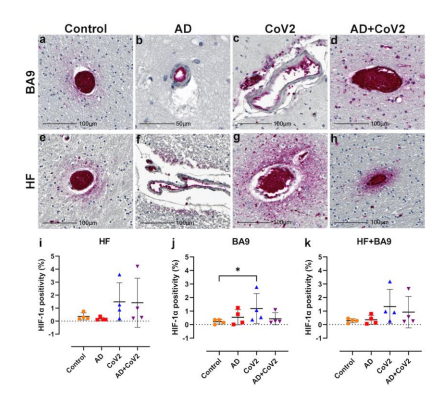




Fig. 6. Hypoxia-inducible factor-1 alpha subunit (HIF-1α) in neurological controls, SARS CoV-2, AD and SARS-CoV-2 infected individuals.

Representative images demonstrate HIF-1 α immunopositivity around the vasculature that extends 714 into the parenchyma in brain of patients with SARS-CoV-2 infection, regardless of AD status 715 shown in panels C, D, G, H. Comparatively, this is seen less frequently and does not extend 716 significantly into the parenchyma in brain of age-matched controls, shown in panels A and E. In 717 AD only, positivity was most often observed in epithelium with no or minimal extension into the 718 719 brain parenchyma as shown in panels **B** and **F**. Graphs show the total percentage of tissue positive for HIF-1a, as defined using a HALO area algorithm for detection of Vector Red 720 intensity over the whole section shown (I-K). Although statistical significance between groups 721 was not reached in the HF or averaged group comparisons, increased HIF-1 α expression is seen 722 in the SARS-CoV-2 infected patients, with and without AD, as compared to age-matched controls 723 in panels I and K. An increase in area positivity is seen in all groups in the cortical BA9 region. 724 as compared to non-affected controls, however, statistical significance is only seen with the 725 726 SARS-CoV-2 group in panel J. Statistics were performed with a two-tailed Mann-Whitney U test. *p<0.05. Data are expressed as mean \pm SEM. 727