

Discovering Common Pathogenic Mechanisms of COVID-19 and Parkinson Disease: An Integrated Bioinformatics Analysis

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Abstract

Coronavirus disease 2019 (COVID-19) has emerged since December 2019 and was later characterized as a pandemic by WHO, imposing a major public health threat globally. Our study aimed to identify common signatures from different biological levels to enlighten the current unclear association between COVID-19 and Parkinson's disease (PD) as a number of possible links, and hypotheses were reported in the literature. We have analyzed transcriptome data from peripheral blood mononuclear cells (PBMCs) of both COVID-19 and PD patients, resulting in a total of 81 common differentially expressed genes (DEGs). The functional enrichment analysis of common DEGs are mostly involved in the complement system, type II interferon gamma (IFNG) signaling pathway, oxidative damage, microglia pathogen phagocytosis pathway, and GABAergic synapse. The protein–protein interaction network (PPIN) construction was carried out followed by hub detection, revealing 10 hub genes (MX1, IF127, C1QC, C1QA, IF16, NFIX, C1S, XAF1, IF135, and ELANE). Some of the hub genes were associated with molecular mechanisms such as Lewy bodies–induced inflammation, microglia activation, and cytokine storm. We investigated regulatory elements of hub genes at transcription factor and miRNA levels. The major transcription factors regulating hub genes are SOX2, XAF1, RUNX1, MITF, and SPI1. We propose that these events may have important roles in the onset or progression of PD. To sum up, our analysis describes possible mechanisms linking COVID-19 and PD, elucidating some unknown clues in between.

Keywords COVID-19 · Parkinson disease · Transcriptome analysis · Regulatory networks · Signaling pathways · Bioinformatics

Abbreviations		CS	Cytokine storm
BP	Biological process	ETS	E26 transformation-specific
CC	Cellular component	GABA	Gamma-aminobutyric acid
CNS	Central nervous system	GEO	Gene Expression Omnibus
COVID-19	Coronavirus disease 2019	GO	Gene Ontology
		HCV	Hepatitis C virus
Aria Jahanimoghadam and Hadis Abdolahzadeh equally contributed to this work.		IFI6	Interferon alpha-inducible protein 6
		IFI27	Interferon alpha-inducible protein 27
		IFI35	Interferon alpha-inducible protein 35
☐ Javad Zahiri jzahiri@health.ucsd.edu		IFN	Interferon
		KEGG	Kyoto Encyclopedia of Genes and
Bioinformatics and Computational Omics Lab (BioCOOL), Department of Biophysics, Faculty of Biological Sciences, Tarbiat Modares University, Tehran, Iran			Genomes
		Lrrk2	Leucine-rich repeat kinase 2
		MF	Molecular functions
Department of Stem Cells and Developmental Biology,		miRNA	microRNAs
	Research Center, Royan Institute for Stem Cell	MITF	Microphthalmia-associated transcription
	Technology, ACECR, Tehran, Iran		factor
	of Neuroscience, University of California San la, San Diego, CA, USA	MX	Myxovirus resistance genes
4		MX1	MX dynamin-like GTPase 1
·	ılius-Maximilians-Universität Würzburg, Am ürzburg, Germany	PBMCs	Peripheral blood mononuclear cells



PD Parkinson's disease

PPIN Protein-protein interaction network

RIN3 Rab interactor 3

RUNX1 Runt-related transcription factor 1

SARS-CoV-2 Severe acute respiratory syndrome corona-

virus 2

SOX2 Sex-determining region Y-box 2

TFs Transcription factors

UCHL1 Ubiquitin carboxyl-terminal hydrolase L1 XAF1 X-linked inhibitor of apoptosis-associated

factor 1

Introduction

According to the statistics provided by WHO, there have been approximately 255 million cases diagnosed with coronavirus disease 2019 (COVID-19) with more than 5 million confirmed death cases as of November 2021. The novel severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) causes COVID-19 and is considered a worldwide pandemic whose impacts are noticeable across the globe (Chams et al. 2020; Khorsand et al. 2020). COVID-19 is initially viewed as a respiratory disease. However, it was demonstrated that SARS-CoV-2 affects multiple organs such as the central nervous system (CNS). Moreover, neurological manifestation associated with SARS-CoV-2 can potentially occur via direct invasion of the virus into the CNS and, thus, introduce the brain as a location containing high replicative values for SARS-CoV-2 (Song et al. 2021; Szcześniak et al. 2021).

Parkinson's disease (PD) is a neurodegenerative disease characterized for the first time in 1817 by James Parkinson. However, after nearly two centuries of research, the cause of most of the cases is still unclear (Olsen et al. 2018; Hayes 2019). There are multiple reports regarding the occurrence of neurological complications in individuals with COVID-19 as up to 85%. Additionally, hyposmia, one of the symptoms of PD, has been reported to happen in 65% of cases with COVID-19 (Merello et al. 2021). The prevalence of PD among elderly ages higher than 65 is 1–3%, and that for the whole population is approximately 0.3%. Of note, PD is distinguished mainly via degeneration of dopaminergic neurons located in the substantia nigra of the midbrain (Kalia and Lang 2016). Subsequently, a number of theoretical mechanisms such as basal ganglia injury, microglia-induced inflammation, and post-encephalopathy inflammation have been proposed to be the hypothetical link between COVID-19 and PD but there is still a gap of knowledge in the way of understanding the relationship among them. This hypothesis is also supported by some case reports, stating a rapid onset of PD after infection with SARS-CoV-2 (Eichel et al. 2020; Cartella et al. 2021; Merello et al. 2021).

On the other hand, several viruses including Epstein-Barr, hepatitis C, herpes simplex 1, influenza A, and varicella-zoster have previously been shown to be related to increasing the risk of diagnosing with PD in the distant future (Henry et al. 2010; Merello et al. 2021). These viruses can directly induce neuronal injury after the infection. For instance, it has been demonstrated that there is a marked increased risk of developing PD after hepatitis C virus (HCV) infection. This was enabled by the ability of HCV to replicate in the CNS (Tsai et al. 2016). The family of Coronaviridae has been known to cause CNS infection (Bergmann et al. 2006), presumably in the case of SARS-CoV-2 via the blood-brain barrier (BBB) due to the cytokine storm (CS) (Eldeeb et al. 2020; Sulzer et al. 2020). Moreover, SARS-CoV-2 signature was detected in the autoptic brains of 21 out of 40 patients (53%) after dying of COVID-19. Although no relationship between the presence of SARS-CoV-2 and the severity of the disease was found, it was proved that SARS-CoV-2 can reach the CNS (Matschke et al. 2020). Overall, all the aforementioned pieces of evidence create an urgent need that the possible crosstalk between SARS-CoV-2 and neurodegenerative disorders such as PD should be taken into consideration. Other studies have been carried out assessing other potential comorbidities in respect to COVID-19 including chronic kidney disease (Auwul et al. 2021) and diabetes mellitus (Rahman et al. 2021).

In the present study, we adopted an integrated bioinformatics analysis to scrutinize the common molecular mechanisms involved in COVID-19 and PD pathogenesis and how SARS-CoV-2 can possibly contribute to developing PD whether immediately after contracting COVID-19 or years later (an overview of the present study is shown in Fig. 1).

Materials and Methods

Transcriptomic Data Analysis

We obtained the transcriptome data from the Gene Expression Omnibus (GEO; https://www.ncbi.nlm.nih.gov/geo/) with accession numbers GSE152418 (16 peripheral blood mononuclear cell (PBMC) samples from COVID-19 subjects and 17 from normal individuals) (Arunachalam et al. 2020) and GSE165082 (12 PBMC samples from PD and 14 from normal individuals) (Henderson et al. 2021). The R package DESeq2 was provided for normalization and differential expression analysis (Love et al. 2014). We used the *P* value < 0.05 and (log FC > | 11) as thresholds. Common differentially expressed genes (DEGs) between two datasets were obtained using the Venny 2.1.0 tool (https://bioinfogp.cnb.csic.es/tools/venny/index.html).



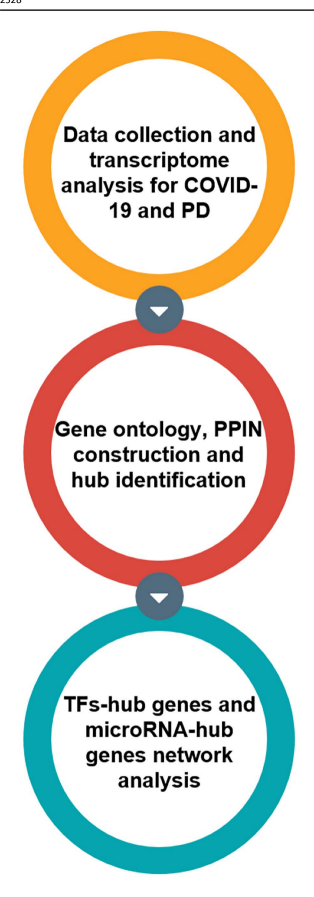


Fig. 1 Flow chart of steps conducted in the study



Gene Ontology and Pathway Enrichment Analysis

For the functional annotation and pathway enrichment analysis of the DEGs, Enrichr web utility tools (Kuleshov et al. 2016) were used. WikiPathways and the Kyoto Encyclopedia of Genes and Genomes (KEGG) were used for finding pathway enrichment analysis. Gene Ontology (GO) terms were considered in three main categories such as biological process (BP), cellular component (CC), and molecular functions (MF).

Protein-Protein Interaction Network Construction and Analysis

GeneMANIA (Warde-Farley et al. 2010) server was used for protein–protein interaction network (PPIN) construction, and then the obtained PPIN was analyzed and visualized by Cytoscape version 3.8. We adopted a hub detection approach called maximal clique centrality (MCC) via cytoHubba plug-in of Cytoscape to retrieve the top 10 hub nodes. MCC is a local-based algorithm which outperforms other methods in hub identification (Chin et al. 2014).

Identification of Transcription Factors and MicroRNAs Regulating Hub Genes

Transcription factor (TF) and microRNA (miRNA) are considered the major regulatory elements of gene expression at both transcription and post-transcription levels (Qin et al. 2020). We have constructed TF-hub gene and miRNA-hub gene regulatory networks with the use of NetworkAnalyst 3.0 to detect important regulatory elements (Zhou et al. 2019). We used ChEA as a TF database (Lachmann et al. 2010) to create the TF-hub gene interaction network. To construct the miRNA-hub gene interaction network, TarBase (Karagkouni et al. 2018) was selected to retrieve interacting miRNAs with regard to hub genes. Following the network's construction, network analysis was carried out to identify core TFs and miRNAs based on the degree.

Results

Identification of Common DEGs Between COVID-19 and PD

We examined transcriptional signatures between COVID-19 (n=16) and healthy controls (n=17). There were 4795 DEGs in COVID-19 versus healthy controls. Also, we obtained DEGs between PD (n=14) and normal subjects (n=12). Our results showed 233 DEGs in PD compared to

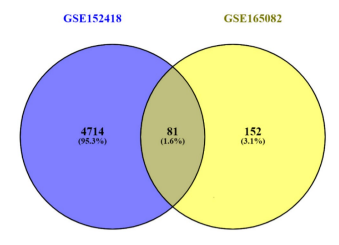


Fig. 2 Venn diagram showing common DGEs between COVID-19 and PD

controls. We detected 81 common DEGs between COVID-19 and PD (Fig. 2). Top ten common DEGs are shown in Table 1.

Pathway Enrichment Analysis

Functional annotations of common DEGs indicated involvement in multiple pathways including the complement system, type II interferon gamma (IFNG) signaling pathway, oxidative damage, microglia pathogen phagocytosis pathway, and GABAergic synapse (Table 2). The GO analysis of common DEGs revealed that enriched BPs were mostly involved in the regulation of complement activation, regulation of immune effector process, regulation of humoral immune response, cell junction disassembly, commissural neuron axon guidance, synapse pruning, complement activation, classical pathway, determination of left/right symmetry,

Table 1 Top ten common DEGs between COVID-19 and PD

The common DEGs	log FC			
	GSE152418 (COVID- 19 versus healthy control)	GSE165082 (Parkinson versus healthy control)		
ISG15	1.2	-1.3		
GABRD	1.8	-2.1		
C1QC	3.5	-1.05		
C1QB	2.9	-1.2		
IFI6	2.05	-1.2		
A3GALT2	2.08	1.04		
VCAM1	-1.2	2.9		
AQP10	1.9	-1.2		
ACTG1P25	1.1	-1.2		
C4BPA	1.1	-1.4		

negative regulation of humoral immune response mediated by circulating immunoglobulin, and humoral immune response mediated by circulating immunoglobulin. The enriched molecular functions were involved in kinase activator activity, protein kinase activator activity, GABA-A receptor activity, neurotransmitter receptor activity involved in the regulation of postsynaptic membrane potential, GABA receptor activity, transmitter-gated ion channel activity involved in the regulation of postsynaptic membrane potential, protein kinase regulator activity, transmitter-gated ion channel activity, glycerol channel activity, and arylesterase activity. CC enriched in GABA-A receptor complex, azurophilic granule, collagen-containing extracellular matrix, Golgi lumen, secretory granule lumen, caveola, specific granule lumen, vacuolar lumen, primary lysosome, and plasma membrane raft (Table 3).

Protein-Protein Interaction Network Construction and Analysis

We constructed PPIN, containing 71 nodes and 1369 edges as shown in Fig. 3. The PPIN depicts the interaction of common DEGs and was visualized by Cytoscape software. According to Table 4, the ten hub genes based on MCC score are myxovirus resistance genes (MX) dynamin-like GTPase 1 (MX1), interferon alpha-inducible protein 27 (IFI27), C1CQ, C1QA, interferon alpha-inducible protein 6 (IFI6), NFIX, C1S, X-linked inhibitor of apoptosis-associated factor-1 (XAF1), interferon alpha-inducible protein (IFI35), and elastase, neutrophil expressed (ELANE). These hub genes can potentially be used as drug targets and play a crucial role in maintaining the stability of the network. Therefore, further analysis of these genes is of great importance. For instance, scrutinizing the regulatory interaction of hub genes is recommended.

Regulatory Networks

In order to gain deeper insights into our hub genes, we sought to construct TF-hub gene and miRNA-hub gene networks. Figures 4 and 5 display the regulators of the hub genes (TFs and miRNAs, respectively). From these regulatory networks, it can be concluded that some regulatory elements are more important and can subsequently interact with more hub genes. In the TF-hub gene network, 16 TFs were identified with 3 or more interactions, whereas in miRNA-hub genes, 29 miRNAs were detected with at least 3 or more interactions. The most connected TFs were SOX2 and XAF1 with the degree of 6, and RUNX1, MITF, SPI1, and MYC with 5 interactions. The most significant miRNA related to hub genes is hsa-mir-129–2-3p with a degree of 8.



Table 2 Top ten molecular pathways enriched by 81 common DEGs in COVID-19 and PD

Source	Pathways	P value	Count	Genes
WikiPathways	Complement and coagulation cascades (WP558)	0.001682	3	C1QB;SERPING1;C1QC
	Complement activation (WP545)	0.003550	2	C1QB;C1QC
	Complement system (WP2806)	0.007179	3	SERPING1;C4BPA;ELANE
	Type II interferon gamma (IFNG) signaling (WP619)	0.009842	2	IFI6;ISG15
	miRNAs' involvement in the immune response in sepsis (WP4329)	0.009842	2	VCAM1;ELANE
	Oxidative damage (WP3941)	0.011437	2	C1QB;C1QC
	Microglia pathogen phagocytosis pathway (WP3937)	0.011437	2	C1QB;C1QC
	Development of ureteric collection system (WP5053)	0.015565	2	WNT11;SMO
	Prader-Willi and Angelman syndrome (WP3998)	0.025409	2	GABRR2;GABRD
	Non-genomic actions of 1,25-dihydroxyvitamin D ₃ (WP4341)	0.033623	2	RSAD2;ISG15
KEGG	Pertussis	0.000256	4	C1QB;SERPING1;C4BPA;C1QC
	Complement and coagulation cascades	0.000394	4	C1QB;SERPING1;C4BPA;C1QC
	Systemic lupus erythematosus	0.002210	4	C1QB;CTSG;ELANE;C1QC
	Neuroactive ligand-receptor	0.002609	6	GABRR2;CHRND;GRID1;LPAR 1;CTSG;GABRD
	Staphylococcus aureus infection	0.006778	3	C1QB;DEFA4;C1QC
	Transcriptional misregulation	0.007708	4	ETV7;DEFA4;ERG;ELANE
	Nicotine addiction	0.011437	2	GABRR2;GABRD
	Basal cell carcinoma	0.026977	2	WNT11;SMO
	GABAergic synapse	0.050585	2	GABRR2;GABRD
	Morphine addiction	0.052626	2	GABRR2;GABRD

Other major miRNAs are hsa-mir-124-3p, hsa-mir-34a-5p, hsa-mir-21-3p, and hsa-mir-27a-5p; each has 6 edges with hub genes.

Discussion

The COVID-19 outbreak has undoubtedly become an international concern (WHO 2021). Some case reports hypothesized rapid onset of PD happens after SARS-CoV-2 infection (Cartella et al. 2021; Merello et al. 2021). However, there is no study aimed to investigate common links between COVID-19 and PD yet in an in silico manner.

In this study, we adopted a network-based approach following transcriptome analysis to detect the common molecular pathways involved in COVID-19 and PD pathogenesis. The analysis demonstrated 81 common DEGs between COVID-19 and PD. We then performed the pathway enrichment analysis of common DEG. Our results showed the complement and coagulation cascades are one of the pathways that are enriched by the common DEGs. The complement system plays a double role in the immune response against SARS-CoV-2 and the pathogenesis of COVID-19 tissue involvement (Gao et al. 2020; Diao et al. 2021). Several studies reported complement components to alter within the blood of PD patients (Goldknopf et al. 2006). The type II IFNG signaling pathway was also identified.

The interferon (IFN) responses constitute the main first line of defense against SARS-CoV-2 (Park and Iwasaki 2020). IFN-y has a role in inflammation and neurodegeneration in PD, as an increase of IFN-γ was detected in the serum of PD patients (Baba et al. 2005). Another common pathway was oxidative damage. Oxidative stress most likely impacts COVID-19 pathogenesis by accompanying cell activation (Chernyak et al. 2020). Oxidative stress is one of the mechanisms mentioned in the etiopathogenesis of PD (Dorszewska et al. 2021). Oxidative stress causes damage to key cellular components in the substantia nigra (SN) of PD patients (Dias et al. 2013). We detected microglia pathogen phagocytosis pathway in which microglia by some pathogenic mechanisms could contribute to the development of post-COVID-19 neurological sequelae and disorders, including PD (Awogbindin et al. 2021). Another enriched pathway was GABAergic synapse. COVID-19-associated inflammation may induce a cortical impairment of GABAergic neurotransmission, possibly representing cognitive fatigue, apathy, and executive deficits (Ortelli et al. 2021). GABA has also been reported to be involved in neurodegenerative disorders such as PD (Muñoz et al. 2020).

The hub genes have been identified from the PPIN to detect major signaling elements that may be used as therapeutic targets for the development of novel drugs to treat COVID-19 patients with PD comorbidity. MX1 is one of the *MX* which has the antiviral effect against RNA viruses. MX1



Table 3 GO enrichment analysis of 81 common DEGs in COVID-19 and PD

	Term	P value	Count	Genes
BP	Regulation of complement activation (GO:0030449)	0.000049	4	C1QB;SERPING1;C4BPA;C1QC
	Regulation of immune effector process (GO:0002697)	0.000062	4	C1QB;SERPING1;C4BPA;C1QC
	Regulation of humoral immune response (GO:0002920)	0.000067	4	C1QB;SERPING1;C4BPA;C1QC
	Cell junction disassembly (GO:0150146)	0.00024	2	C1QB;C1QC
	Commissural neuron axon guidance (GO:0,071679)	0.00044	2	SMO;NFIB
	Synapse pruning (GO:0098883)	0.00057	2	C1QB;C1QC
	Complement activation, classical pathway (GO:0006958)	0.00057	2	C1QB;C1QC
	Determination of left/right symmetry (GO:0007368)	0.00065	3	DNAH11;SMO;FOXJ1
	Negative regulation of humoral immune response mediated by circulating immunoglobulin (GO:0002924)	0.00071	2	FOXJ1;C4BPA
	Humoral immune response mediated by circulating immunoglobulin (GO:0002455)	0.00087	2	C1QB;C1QC
MF	Kinase activator activity (GO:0019209)	0.00026	3	WNT11;SPDYA;GPRC5D
	Protein kinase activator activity (GO:0030295)	0.002133	3	WNT11;SPDYA;GPRC5D
	GABA-A receptor activity (GO:0004890)	0.002649	2	GABRR2;GABRD
	Neurotransmitter receptor activity involved in the regulation of postsynaptic membrane potential (GO:0099529)	0.003236	2	CHRND;GRID1
	GABA receptor activity (GO:0016917)	0.00355	2	GABRR2;GABRD
	Transmitter-gated ion channel activity involved in the regulation of postsynaptic membrane potential (GO:1904315)	0.006547	2	CHRND;GRID1
	Protein kinase regulator activity (GO:0019887)	0.007384	3	WNT11;SPDYA;GPRC5D
	Transmitter-gated ion channel activity (GO:0022824)	0.008356	2	CHRND;GRID1
	Glycerol channel activity (GO:0015254)	0.020088	1	AQP10
	Arylesterase activity (GO:0004064)	0.024058	1	CA1
CC	GABA-A receptor complex (GO:1902711)	0.002649	2	GABRR2;GABRD
	Azurophilic granule (GO:0042582)	0.003635	4	CEACAM6;DEFA4;CTSG;ELANE
	Collagen-containing extracellular matrix (GO:0062023)	0.004433	6	C1QB;SERPING1;CTSG;CSPG4;ELANE;C1QC
	Golgi lumen (GO:0005796)	0.007805	3	DEFA4;CSPG4;MUC5B
	Secretory granule lumen (GO:0034774)	0.009194	5	DEFA4;SELENOP;SERPING1;CTSG;ELANE
	Caveola (GO:0005901)	0.02464	2	SMO;KCNA5
	Specific granule lumen (GO:0035580)	0.026188	2	DEFA4;ELANE
	Vacuolar lumen (GO:0005775)	0.027664	3	CTSG;CSPG4;ELANE
	Primary lysosome (GO:0005766)	0.043669	1	DEFA4
	Plasma membrane raft (GO:0044853)	0.043678	2	SMO;KCNA5

BP biological processes, MF molecular functions, CC cellular components

expression has been reported to be elevated in COVID-19 patients and conversely declines as age increases. Plus, it can be stimulated in the cytoplasm by IFNs and participates in the cellular antiviral response to SARS-COV-2 (Bizzotto et al. 2020). Furthermore, the accumulation of α -synuclein (α -SYN) in the brain of PD patients induces the expression of MX1. This molecule is involved in PI3K-Akt signaling pathway, cytokine release, and immune response IFN- α , IFN- β , and IFN- γ signaling pathways (Yamada et al. 1994; Qin et al. 2016). It is also a regulator of IFN systems that contributes to CS (Yang et al. 2021). This might facilitate the entry of virus to the CNS via the BBB. It is noteworthy that the BBB was reported to be disrupted in the animal

models of PD which can lead to degeneration of neurons in the substantia nigra (Al-Bachari et al. 2020). MX1 localized in self-aggregations and generated Lewy bodies and swelling of neuronal processes in the substantia nigra of brain tissues in Parkinson's patients (McDonough et al. 2017). Lewy bodies which contain misfolded proteins can then trigger the activation of T cells (Sulzer et al. 2017). IFN-alpha inducible (IRI) family members are closely related to the inflammatory immune response in COVID-19 and PD (Shaath et al. 2020). IFI6 is an immune-associated early predictor for PD (Lei et al. 2020; Yu et al. 2020). IFI35 is involved in type I interferon signaling pathway and have a vital role in inflammation response in SARS-CoV-2-infected cells (Hachim



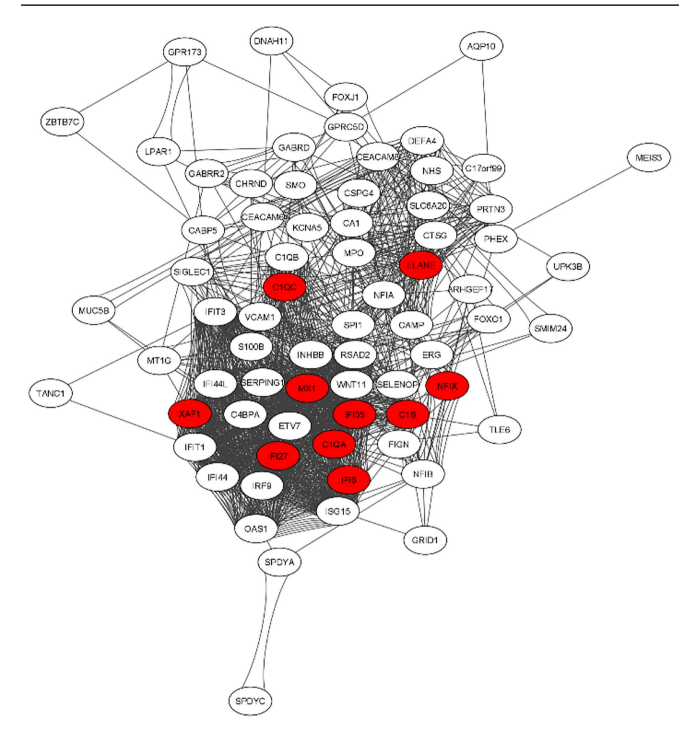


Fig. 3 PPIN of common DEGs. Red nodes indicate top 10 hub genes identified by MCC

et al. 2020; Ziegler et al. 2020; Ong et al. 2021). On the other hand, IFI35 is upregulated in PD patients in response to INF response (Yu et al. 2020). IFI35 gene is expressed in the stratum and substantia nigra regions of the brain, and its de novo mutation is contributed to early onset of PD pathogenesis (Guo et al. 2018). IFI27 is an early predictor for SARS-COV-2 infection, and high-level expression of IF127

is associated with the presence of a high viral load (Shojaei et al. 2021). One study found elevated expression of IFI27 after microglial activation and neuroinflammation in progressive neurodegenerative disorders such as PD (Zhou et al. 2015). SARS-CoV-2 infection induces a strong activation of major constituents of the human complement subcomponent C1q (C1QA, C1QB, C1QC) (Ramlall et al. 2020;



Table 4 Summary of hub nodes

Hubs	MCC
MX1	5
IFI27	4
C1QC	3
C1QA	3
IFI6	3
NFIX	2
C1S	2
XAF1	2
IFI35	2
ELANE	2

MCC maximal clique centrality

Santiesteban-Lores et al. 2021). These genes are upregulated in the microglial cells in the brain of PD patients. Activation of the complement system improves the removal of pathogens and products of tissue damage from the brain and is related to neuronal cell death in PD (Depboylu et al. 2011; Mariani et al. 2016; Itoh and Voskuhl 2017). ELANE gene codes destructive enzymes named neutrophil elastase that play a key role in host defense mechanism. This enzyme

is highly overexpressed in naso-oropharyngeal and blood samples of COVID-19 patients. Neutrophil elastase can activate the spike (S) protein and mediate viral entry and pathogenesis of SARS-COV-2 (Belouzard et al. 2010; Akgun et al. 2020; Guéant et al. 2021). After an inflammatory insult to the CNS structure, the expression of neutrophil elastase increases, then degrades basal lamina and extracellular matrix (ECM) molecules and suppresses neurobehavioral recovery mechanisms (Stowe et al. 2009; Stock et al. 2018). Neutrophil elastase inhibitors could be new treatment options for COVID-19 patients (Mohamed et al. 2020).

Among these transcription factors, sex-determining region Y-box 2 (SOX2) has a critical role in the development and maintenance of neural stem/progenitor cell populations committed to becoming glial cells. SOX2 inhibits myelination in the peripheral nerves and maintains Schwann cells in a proliferative state, which is also associated with the influx of macrophages and increased neuroinflammation (Roberts et al. 2017). Interestingly, the expression level of SOX2 was found to be elevated in the brains of PD patients (Vedam-Mai et al. 2014). Nerve inflammation is one of the important factors in the onset or progression of PD (Pajares et al. 2020). XAF1 is a mitochondrial apoptosis activator

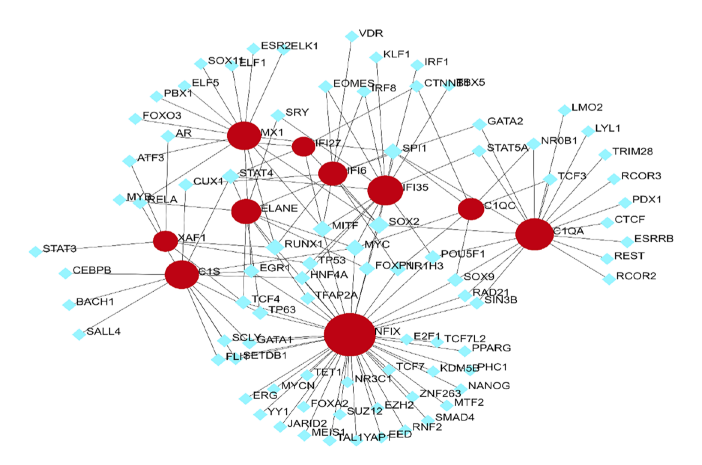
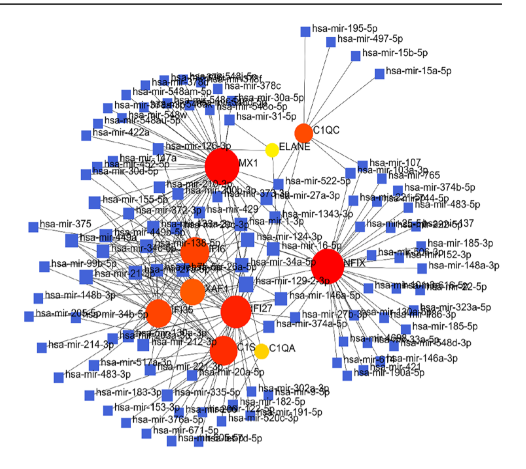


Fig. 4 TF-hub gene regulatory network acquired from Network Analysis web server. Square nodes representing TFs and circle nodes are stand for hub genes



Fig. 5 miRNA-hub gene regulatory network acquired from Network Analysis web server. Square nodes represent miR-NAs that regulate circle nodes which denote hub genes



that is upregulated in immune cells (T, B, natural killer, and dendritic cells) of COVID-19 patients that may be associated with increased apoptosis of these cells (Zhu et al. 2020; Gao et al. 2021). Furthermore, XAF1 expression is higher in the midbrain of PD patients (Gispert et al. 2015; Santiago and Potashkin 2017). IFN-α and IFN-β induced XAF1 mRNA expression and therefore induced cell apoptosis (Leaman et al. 2002). The expression of Runt-related transcription factor 1 (RUNX1) increases after SARS-CoV-2 infection (O'Hare et al. 2021). Interestingly, its overexpression is related to the progression of PD. RUNX1 increases the expression of leucine-rich repeat kinase 2 (*Lrrk2*) gene in immune cells and has a critical role in the pathogenesis of familial PD due to developing hyperactive inflammatory phenotype, neuronal toxicity, and cell apoptosis (Cook et al. 2017; Thomsen et al. 2021). Microphthalmia-associated transcription factor (MITF) is one of the key TFs with varying functions in cell homeostasis, cell cycle, and apoptosis. MITF is upregulated in immune cells and worsens severity of infection in an unknown way in COVID-19 patients (Bost et al. 2020; Ding et al. 2021; Jeong et al. 2021). Ubiquitin carboxyl-terminal hydrolase L1 (UCHL1) is expressed in neural cells and inhibits the stability of MITF by binding to the ubiquitinated protein. The ligase activity of UCHL1 is disrupted in PD, resulting in MITF overexpression and cell

damage in these patients (Liu et al. 2002; Seo et al. 2017). The E26 transformation–specific (ETS) family transcription factor SPI1 upregulated in PBMCs of COVID-19 patients and is involved in the inflammatory process and modulates host immune systems of these patients (Fagone et al. 2020; Rahman et al. 2021). SPI1 plays a key role in the identity, differentiation, and specialized functions of microglia. Microglia rapidly activate in response to pro-inflammatory response. These activated microglia are accumulated in brain lesions of PD patients. SPI1 has many target functional genes in microglial cells including Spi1, Runx1, Irf8, Il34, Aif1, Csf1r, Csf1, Cx3cr1, Tyrobp, and Trem2 (Satoh et al. 2014). SPI1 induces cytokine release and microglial pro-inflammatory response (Pimenova et al. 2021). Therefore, misregulation of SPI1 target genes might lead to the establishment or development of PD due to the accumulation of activated microglia (Satoh et al. 2014). In addition, one multi-omic study identifies a single nucleotide polymorphism, rs10130373, within a microglia-specific peak; interrupts a SPI1 motif; and interfaces effectively with the promoter of the Rab interactor 3 (RIN3) gene. RIN3 plays an important role in the early endocytic pathway that needs microglial function, thereby playing a particularly critical role in progressive neurodegenerative disease (Kajiho et al. 2003; Corces et al. 2020).



hsa-mir-129–2-3p is the most significant miRNA in miRNA-hub gene regulatory networks. miR-129 is a brainenriched miRNA, and its level increases in the peripheral blood lymphocytes of PD patients (Qin et al. 2016).

In the present study, an integrated bioinformatics approach was adopted to explore the possible risk of PD development after COVID-19 infection by investigating the common molecular mechanisms. By taking advantage of the holistic viewpoint of systems biology, we were able to consider every aspect of both diseases and infer novel hypotheses. Further supplementary studies need to be conducted to clarify the association between COVID-19 and PD, as, at the moment, there is little known regarding both of these disease entities. It is worth mentioning that contracting PD is a complex and age-dependent neurodegenerative disorder. Thus, it is encouraged to investigate infected COVID-19 patients' years after their infection to estimate the probability of getting PD.

Conclusion

The current study aimed to investigate common regulators between COVID-19 and PD. Overall, our analysis highlights multiple mechanisms such as complement system, oxidative stress, activation microglia, cytokine storm, and activation of T cells by misfolded proteins which might be the potential links between both comorbidities. Nonetheless, as this is a thorough in silico analysis, the results of this work should be taken into account carefully. Further case reports and follow-up experiments of COVID-19 patients can corroborate these links.

Author Contribution All authors contributed to the study conception and design. Material preparation, data collection, and analysis were performed by *Aria Jahanimoghadam* and *Hadis Abdolahzade*. The first draft of the manuscript was written by *Aria Jahanimoghadam*, *Hadis Abdolahzadeh*, and *Niloofar Khoshdel rad*, and all authors commented on previous versions of the manuscript. All authors read and approved the final manuscript.

Availability of Data and Materials The data used in this study were downloaded from the GEO database.

Code Availability The code that supports the findings of this study is available on request from the corresponding author.

Declarations

Ethics Approval and Consent to Participate Not applicable.

Consent for Publication Not applicable.

Competing Interests The authors declare no competing interests.

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