Review

Blood and Cerebrospinal Fluid Biomarkers of Inflammation in Parkinson's Disease

Milan Zimmermann^{a,b} and Kathrin Brockmann^{a,b,*}

^aCenter of Neurology, Department of Neurodegeneration and Hertie Institute for Clinical Brain Research, University of Tuebingen, Tuebingen, Germany ^bGerman Center for Neurodegenerative Diseases, University of Tuebingen, Tuebingen, Germany

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Abstract. Given the clear role of inflammation in the pathogenesis of Parkinson's disease (PD) and its impact on incidence and phenotypical characteristics, this review provides an overview with focus on inflammatory biofluid markers in blood and cerebrospinal fluid (CSF) in PD patient cohorts. In preparation for clinical trials targeting the immune system, we specifically address the following questions: 1) What evidence do we have for pro-inflammatory profiles in blood and in CSF of sporadic and genetic PD patients? 2) Is there a role of anti-inflammatory mediators in blood/CSF? 3) Do inflammatory profiles in blood reflect those in CSF indicative of a cross-talk between periphery and brain? 4) Do blood/CSF inflammatory profiles associated with phenotypical trajectories in PD? 6) Are blood/CSF inflammatory profiles associated with CSF levels of neurodegenerative/PD-specific biomarkers? Knowledge on these questions will inform future strategies for patient stratification and cohort enrichment as well as suitable outcome measures for clinical trials.

Keywords: Parkinson's disease, inflammation, interleukin, chemokine, cytokine, blood, cerebrospinal fluid

ABBREVIATIONS:

Alpha Fetoprotein	AFP
Anti-Neutrophil Activating	NAP-3 (CXCL1)
Protein 3	
Brain Derived Neurotrophic	BDNF
Factor	
Complement component 3	C3
Complement component 4	C4
Cancer Antigen 125	CA-125

*Correspondence to: Kathrin Brockmann, MD, Department of Neurodegeneration, Hertie Institute for Clinical Brain Research, German Center for Neurodegenerative Diseases, University of Tuebingen, Hoppe Seyler-Strasse 3, 72076 Tuebingen, Germany. Tel.: +49 7071 2980171; Fax: +49 7071 294617; E-mail: kathrin. brockmann@uni-tuebingen.de.

Carcinoembryonic Antigen	CEA			
C-reactive Protein	CRP			
Creatine Kinase Muscle	CKMB			
Brain type				
Eotaxin	CCL11			
Epithelial-derived neutrophil-	ENA-78 (CXCL5)			
activating peptide 78				
FactorVII	FactorVII			
Fatty Acid Binding Protein	FABP			
Fractalkine	CX3CL1			
Growth Hormone	GH			
Immunoglobulin E	IgE			
Intercellular Adhesion	ICAM-1			
Molecule 1				
Interferon-Gamma	IFN-γ			
Interferon-Gamma Induced	IP-9 (CXCL11)			

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Protein-9 Interferon-Gamma Induced	IP-10 (CXCL10)	Epithelial Chemokine Neutrophil Gelatinase-	NGAL
Protein-10	II -10 (CACL10)	Associated Lipocalin	NOLL
Interleukin-1 alpha	IL-1α	NLR Family Pyrin Domain	NLRP3
Interleukin-1 beta	IL-1β	Containing 3	
Interleukin-1 Receptor	IL-1RA	Platelet-Derived Growth	PDGF-BB
Antagonist		Factor BB	
Interleukin-2	IL-2	Prolactin	Prolactin
Interleukin-3	IL-3	Prostate Specific Antigen	PSA-f
Interleukin-4	IL-4	(free)	
Interleukin-5	IL-5	RANTES	CCL5
Interleukin-6	IL-6	S100 Calcium-Binding	S100A8/9
Interleukin-7	IL-7	Protein A8/9	(Calprotectin)
Interleukin-8	IL-8	S100 Calcium-Binding	S100B
Interleukin-9	IL-9	Protein B	
Interleukin-10	IL-10	Serum Amyloid A	SAA
Interleukin-12p40	IL-12p40	Soluble Interleukin-2	sIL-2R
Interleukin-12p70	IL-12p70	Receptor	
Interleukin-13	IL-13	Soluble Tumor Necrosis	sTNFR1
Interleukin-15	IL-15	Factor Receptor 1	
Interleukin-16	IL-16	Soluble Tumor Necrosis	sTNFR2
Interleukin-17 A	IL-17A	Factor Receptor 2	
Interleukin-18	IL-18	Soluble Vascular Cell	sVCAM1
Interleukin-21	IL-21	Adhesion Molecule 1	
Interleukin-22	IL-22	Stem Cell Factor	SCF
Interleukin-27	IL-27	Stromal Cell-Derived	SDF-1
Leptin	Leptin	Factor-1	
Lymphotactin	Lymphotactin	Thyreoperoxidase	TPO
Macrophage Colony	CSF-1	Thyroid Stimulating	TSH
Stimulating Factor 1		Hormone	
Macrophage Derived	MDC	Tissue Factor	TF
Chemokine		Tissue Inhibitor of	TIMP-2
Macrophage Inflammatory	MIP-1a (CCL3)	Metalloproteinases 2	
Protein 1 Alpha		Transforming Growth	TGF-α
Macrophage Inflammatory	MIP-1β (CCL4)	Factor Alpha	
Protein 1 Beta	• • •	Transforming Growth	TGF-β1
Macrophage Inflammatory	MIP-3α (CCL20)	Factor Beta1	·
Protein 3 Alpha		Tumor Necrosis	TNF-α
Macrophage Inflammatory	MIP-3 (CCL23)	Factor alpha	
Protein 3		Tumor Necrosis	TNF-β
Matrix Metallopeptidase 3	MMP3	Factor beta	·
Matrix Metallopeptidase 9	MMP9	Chitinase-3-like 1	YKL-40
Matrix Metallopeptidase 9	MMP10		
Monocyte Chemoattractant	MCP-1(CCL2)		
Protein 1		INTRODUCTION	
Monocyte Chemoattractant	MCP-2 (CCL8)		
Protein 2		In recent years, a growing	g number of epider
Monocyte Chemoattractant	MCP-4 (CCL13)	logical and genetic studies a	
Protein 4		and biofluid marker analyses	-
Monokine Induced by	MIG (CXCL9)	teomic approaches provide e	
Interferon-Gamma		influence of inflammation on	
Musses Associated	MEC(CCI 28)		

MEC (CCL28)

Mucosae Associated

emioortem d proevant influence of inflammation on both incidence and progression in Parkinson's disease (PD) [1-8]. In this

context, the cerebral and peripheral as well as the innate and adaptive immune system seem involved [9]. The activation of microglia as representative of the cerebral innate immune system was shown postmortem and in vivo in PD patients by positron emission tomography (PET) studies ([¹¹C]PK11195, [¹⁸F]FEPPA, [¹¹C]PBR28) and by increased levels of cytokines in cerebrospinal fluid (CSF) [2, 10]. Microglia is activated by damage-associated molecular patterns (DAMPs), which are generated by damaged cells, misfolded proteins, and protein aggregates. In PD, α -synuclein acts as DAMP resulting in microglia activation with induction of neuroinflammation and release of cytokines/chemokines [11]. Moreover, there is increasing evidence for the involvement of the peripheral innate and adaptive immune system in the pathophysiology of PD. In this context, a-synuclein promotes inflammasomerelated cytokine production in the periphery and specific α -synuclein peptides act as antigenic epitopes resulting in helper and cytotoxic T cell responses in peripheral blood mononuclear cells from patients with PD [12, 13].

Postmortem and biofluid (blood, CSF) studies reported that increased inflammatory profiles are associated with clinical subtypes of PD, promoting an accelerated motor and non-motor phenotype [3, 14–17]. Recent evidence highlights that the involvement of inflammation in PD is maximized in the early disease stages and maintains a chronic profile during the course of the disease [18, 19] (Fig. 1).

Despite this clear role for inflammation in the pathogenesis of PD, several open questions remain to be answered in preparation of clinical trials aiming at disease-modification by targeting the immune system: 1) What evidence do we have for pro-inflammatory profiles in blood and in CSF of sporadic and genetic PD patients? 2) Is there a role of anti-inflammatory mediators in blood/CSF? 3) Do inflammatory profiles in blood reflect those in CSF indicative of a cross-talk between periphery and brain? 4) Do blood/CSF inflammatory profiles change over the disease course as assessed in repeatedly taken biosamples? 5) Are blood/CSF inflammatory profiles associated with phenotypical trajectories in PD? 6) Are blood/CSF inflammatory profiles associated with CSF levels of neurodegenerative/PD-specific biomarkers? 7) Which inflammatory markers in blood/CSF are the most promising candidates for clinical trials with regard to patient stratification, cohort enrichment and outcome measures?

This review gives an overview of the current knowledge on these questions with specific focus on blood (plasma, serum) and CSF levels of inflammatory biofluid markers in PD patient cohorts. We did a PubMed search using the search terms "Parkinson, Inflammation, Interleukin, Chemokine, Cytokine, Blood, CSF" and included publications between 01/2011-01/2022. Studies on white blood cells of the myeloid and lymphoid cell lineage as well as on postmortem tissue, imaging, and cell/animal models are not included in the present manuscript as these are reviewed in other articles from this special issue on "The immune system in Parkinson's disease".

Please note that the assessed cohorts largely vary in sample size and disease duration. Further, the type of assays used, differs and ranges from singleplex (ELISA) to multiplex immunoassays with different metrics of quality control (lower limit of quantification, number of replicates, variation coefficient, etc.). Additionally, it is well known that females and males differ in their immune system. They show distinct patterns in innate and adaptive immune responses which further change across the lifespan [20]. In general, females show stronger innate and adaptive immune responses than males with increased production of antibodies and anti-inflammatory species such as IL-4 and IL-10. Taking these gender differences into consideration is of importance when analysing immune marker profiles in different cohorts against disease status. However, only few studies stratified their analyses by sex. All these points pose limitations and have to be kept in mind for data interpretation and drawing congruent conclusions.

WHAT EVIDENCE DO WE HAVE FOR PRO-INFLAMMATORY PROFILES IN BLOOD AND IN CSF OF SPORADIC AND GENETIC PD PATIENTS?

In preparation of upcoming clinical trials targeting the immune system, it will be essential to translate basic research findings from cell and animal models into patient cohorts and to define markers that are representative of inflammation, that are suited for patient stratification and cohort enrichment, that are predictive/prognostic for different trajectories and that serve as read-out for target engagement. Moreover, such markers should be longitudinally accessible in a multi-centre design. While specific imaging techniques such as PET might not be readily available all

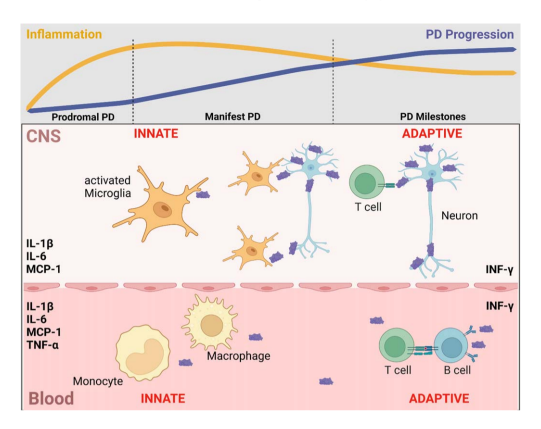


Fig. 1. Inflammatory markers in blood and CSF that are associated with clinical trajectories, disease progression, and neurodegenerative/disease-specific protein levels. In line with evidence from post-mortem and cell studies, blood and CSF levels of inflammatory markers indicate a contribution of the innate and adaptive immune system in both CNS and periphery. Of these, the most frequently reported inflammatory markers in blood and CSF that are associated with clinical trajectories, disease progression and neurodegenerative/disease-specific protein levels are IL-1 β , IL-6, INF- γ , MCP-1 and TNF- α . Notably, inflammation is maximized in the early disease stages and maintains a chronic profile during the course of the disease. Figure created in BioRender.com.

over the world, blood can be repeatedly collected and stored for longitudinal biofluid marker analyses.

Most studies on inflammatory biofluid markers have been conducted cross-sectionally in blood comparing levels of cytokines between patients with sporadic PD and healthy controls.

Inflammatory markers in blood

Over 50 different pro-inflammatory biofluid markers have been assessed in serum or plasma. However, only for 7 markers (CRP, IL-1 β , IL-2, IL-6, IL-8, IFN- γ , TNF- α) more than 5 studies are available. The most robust data are published for CRP [21–34] and IL-1 β [25–27, 35–44]. These two show consistently higher levels in sporadic PD patients compared to healthy controls. Data for IL-2 [25, 27–29, 40, 45, 46], IL-6 [21, 25–29, 35, 37, 39–42, 45–50], IL-8 [25, 28, 29, 35, 46, 51, 52], IFN- γ [25, 26, 28, 29, 40, 46, 50, 53–55], and TNF- α [25–29, 35, 36, 39–41, 46, 50, 51, 53, 54, 56–59] are less clear with some

studies reporting higher blood levels but others showing no differences or even lower levels in sporadic PD patients when compared to healthy controls. A detailed overview of all assessed blood markers along with the respective findings and references is given in Table 1.

Inflammatory markers in CSF

Studies in CSF are less frequent. Overall, 26 proinflammatory markers have been assessed. However, only IL-1 β , IL-6, IL-8, and TNF- α have been measured in 4 or more studies. Of these, levels of IL-1 β were consistently higher in sporadic PD patients compared to healthy controls [36, 37, 60, 61] while levels of IL-6 were also mostly higher in sporadic PD patients but with less robustness across studies [37, 42, 46, 48, 60–63]. Data for IL-8 and TNF- α are less clear with some studies reporting higher CSF levels but others showing no differences or even lower levels in sporadic PD patients when compared

	and CSF compared to healthy controls from 01/201	12-12/2021
	Blood (PD vs. CON)	CSF (PD vs. CON)
C3	Sun et al. [96] 2019 ↓	
C4	Sun et al. [96] 2019 ↓	
CCL2 (MCP-1)	Csencsits-Smith et al. [58] 2016 ↑	Hall et al. [63] 2018 \longrightarrow
	Schröder et al. [46] $2018 \rightarrow$	Schröder et al. [46] 2018 ↑
	Usenko et al. [55] 2020 ↑	
	Miliukhina et al. [50] 2020 \downarrow	
CCL3 (MIP-1a)	Schröder et al. [46] $2018 \rightarrow$	Schröder et al. [46] $2018 \rightarrow$
	Calvani et al. [52] 2020 ↓	
CCL4 (MIP-1β)	Schröder et al. [46] $2018 \rightarrow$	
	Calvani et al. [52] 2020 ↑	
CCL5 (RANTES)	Mahlknecht et al. [45] $2012 \rightarrow$	
	Tang et al. [47] 2014 ↑	
	Qin et al. [25] 2016 ↑	
	Schröder et al. [46] $2018 \longrightarrow$	
CCL8 (MCP-2)		Santaella et al. [65] 2020 ↓
CCL11 (Eotaxin)	Schröder et al. [46] 2018 \rightarrow	
CCL13 (MCP-4)	Mahlknecht et al. [45] $2012 \rightarrow$	
CCL17 (TARC)	Schröder et al. [46] 2018 \rightarrow	
CCL20 (MIP- 3α)	Schröder et al. [46] 2018 \longrightarrow	
CCL23 (MIP-3)		Santaella et al. [65] 2020 \downarrow
CCL28 (MEC)		Santaella et al. [65] 2020 \uparrow
CRP	Ton et al. [21] 2012 \downarrow	Hall et al. [63] 2018 \rightarrow
	Andican et al. [22] 2012 \uparrow	Moghaddam et al. [87] 2018 ↑
	Lindqvist et al. [23] 2013 \uparrow	
	Sawada et al. [24] 2014 \uparrow	
	Qin et al. [25] 2016 \uparrow Wang at al. [26] 2016 \uparrow	
	Wang et al. [26] 2016 ↑ Kim et al. [27] 2018	
	Kim et al. [27] 2018 \longrightarrow King et al. [28, 29] 2019 \longrightarrow	
	Santos-García et al. [30] 2019 ↑	
	Qiu et al. [31] 2019 ↑	
	Baran et al. [32] 2019 \uparrow	
	Jin et al. [33] 2020 \uparrow	
	Dommershuijsen et al. [34] $2022 \rightarrow$	
CSF-1		Santaella et al. [65] $2020 \longrightarrow$
CX3CL1 (Fractalkine)	Gupta et al. [78] 2021 ↑	Santaella et al. [65] 2020 \downarrow
		Hatcher-Martin et al. [89] 2021 \downarrow
CXCL1 (NAP-3)	Schröder et al. [46] 2018 \longrightarrow	Santaella et al. [65] 2020 \downarrow
CXCL5 (ENA78)	Schröder et al. [46] 2018 \rightarrow	
CXCL9 (MIG)	Schröder et al. [46] 2018 \longrightarrow	
CXCL10 (IP-10)	Schröder et al. [46] 2018 \rightarrow	Schröder et al. [46] 2018 \longrightarrow
chello (il 10)	Csencsits-Smith et al. [58] 2016 ↑	Hu et al. [64] 2019 \rightarrow
CXCL11 (IP-9)	Schröder et al. [46] 2018 \rightarrow	110 et al. [04] 2017
CXCL12 (SDF-1)	Bagheri et al. [97] 2018 ↑	
FABP	Brockmann et al. [38] 2016 \uparrow	
ICAM-1	Andican et al. [22] 2012 \uparrow	
	Mahlknecht et al. [45] $2012 \rightarrow$	
IL-1B	Koziorowski et al. [35] $2012 \rightarrow$	Milyukhina et al. [37] 2015 ↑
	Hu et al. [36] 2015 \uparrow	Hu et al. [36] 2015 \uparrow
	Milyukhina et al. [37] 2015 ↑	Chen et al. [60] 2018 \uparrow
	Brockmann et al. [38] 2016 \uparrow	Iwaoka et al. [61] 2020 \uparrow
	Wang et al. [26] 2016 \uparrow	1
	Qin et al. [25] 2016 \uparrow	
	Karpenko et al. [39] 2018 ↑	
	Kin et al. [27] 2018 \uparrow	
	Rocha et al. [40] 2018 \rightarrow	
	Alrafiah et al. [41] 2019 \uparrow	
	Lian et al. [42] 2019 ↑	
	Chatterjee et al. [43] 2020 ↑	
	Fan et al. [44] 2020 ↑	

 Table 1

 Findings of levels of inflammatory markers in sporadic PD patients in blood (serum, plasma) and CSF compared to healthy controls from 01/2012-12/2021

	(Continued)	
	Blood (PD vs. CON)	CSF (PD vs. CON)
П 1D А		
IL-1RA IL-2	Karpenko et al. [39] 2018 \downarrow Mahlknecht et al. [45] 2012 \longrightarrow	Schröder et al. [46] 2018 ↑
IL-2	Qin et al. [25] 2016 \uparrow	Schloder et al. [40] 2018
	Kim et al. [27] 2018 \uparrow	
	Rocha et al. [40] 2018 \rightarrow	
	Schröder et al. [46] 2018 \rightarrow	
	King et al. [28, 29] 2019 \longrightarrow	
sIL-2R	Wang et al. [26] 2016 \uparrow	
IL-4	Qin et al. [25] 2016 \rightarrow	Schröder et al. [46] 2018 \longrightarrow
	Rocha et al. [40] 2018 ↓	
	Schröder et al. [46] 2018 \longrightarrow	
	King et al. $[28, 29]$ 2019 \longrightarrow	
IL-5	Schröder et al. [46] $2018 \rightarrow$	
IL-6	Ton et al. [21] 2012 ↑	Yu et al. [62] 2014: ↑
	Koziorowski et al. [35] $2012 \rightarrow$	Milyukhina et al. [37] 2015 ↑
	Mahlknecht et al. [45] $2012 \rightarrow$	Delgado-Alvarado et al. [48] $2017 \rightarrow$
	Tang et al. [47] 2014 ↑	Chen et al. [60] 2018 ↑
	Milyukhina et al. [37] 2015 ↑	Hall et al. [63] 2018 \longrightarrow
	Wang et al. [26] 2016 ↑	Schröder et al. [46] 2018 ↑
	Qin et al. [25] 2016 ↑	Lian et al. [42] 2019 ↑
	Delgado-Alvarado et al. [48] $2017 \longrightarrow$	Iwaoka et al. [61] $2020 \rightarrow$
	Karpenko et al. [39] 2018 ↑	
	Kim et al. [27] 2018 ↑	
	Rocha et al. [40] 2018 ↓	
	Schröder et al. [46] 2018 \longrightarrow	
	Alrafiah et al. [41] 2019 \longrightarrow	
	King et al. [28, 29] 2019 ↑	
	Lian et al. [42] 2019 ↑	
	Kwiatek-Majkusiak et al. [49] 2020 ↑	
но	Miliukhina et al. [50] 2020 \downarrow	
IL-8	Koziorowski et al. [35] $2012 \rightarrow$	Hall et al. [63] 2018 ↑
	Gupta et al. [51] 2016 \downarrow	Schröder et al. [46] 2018 \longrightarrow
	Qin et al. [25] 2016 \longrightarrow Schröder et al. [46] 2018 \longrightarrow	Hu et al. [64] $2019 \rightarrow$ Santaella et al. [65] $2020 \downarrow$
	King et al. [28, 29] 2019 \longrightarrow	
	Calvani et al. [52] $2019 \longrightarrow$	
IL-9	Schröder et al. [46] 2018 \rightarrow	Schröder et al. [46] 2018 ↓
	Calvani et al. [52] $2020 \downarrow$	
IL-10	Koziorowski et al. [35] $2012 \rightarrow$	Schröder et al. [46] 2018 \longrightarrow
	Brockmann et al. [38] 2016 \uparrow	Hu et al. [64] 2019 \rightarrow
	Qin et al. [25] 2016 \uparrow	
	Karpenko et al. [39] 2018 \rightarrow	
	Kim et al. [27] 2018 \rightarrow	
	Rocha et al. [40] 2018 \downarrow	
	Schröder et al. [46] 2018 \rightarrow	
	King et al. $[28, 29]$ 2019 \longrightarrow	
	Rathnayake et al. [54] 2019 ↑	
	Martin-Ruiz et al. [57] 2020 ↑	
IL-12	Koziorowski et al. [35] 2012 \longrightarrow	
IL-12-p40	Brockmann et al. [38] 2016 ↑	
IL-13	Schröder et al. [46] 2018 \longrightarrow	
	Lin et al. [53] 2019 ↑	
IL-16	·	
IL-17A	Rocha et al. [40] 2018 ↓	
Schröder et al. [46] 2018 \longrightarrow	Schröder et al. [46] 2018 \longrightarrow	Majbour et al. [68] 2020 ↓
IL-21	Schröder et al. [46] 2018 \longrightarrow	Schröder et al. [46] 2018 \longrightarrow
IL-22	Schröder et al. [46] 2018 \longrightarrow	Schröder et al. [46] 2018 \longrightarrow
IL-27	Kouchaki et al. [59] 2018 ↓	

Table 1

(Continued)

	(Continued)	
	Blood (PD vs. CON)	CSF (PD vs. CON)
IFN-y	Wang et al. [26] $2016 \longrightarrow$	Schröder et al. [46] 2018 \longrightarrow
	Qin et al. [25] 2016 \longrightarrow	Iwaoka et al. [61] $2020 \longrightarrow$
	Eidson et al. [56] 2017 ↑	
	Rocha et al. [40] 2018 ↓	
	Schröder et al. [46] 2018 \longrightarrow	
	King et al. [28, 29] $2019 \longrightarrow$	
	Lin et al. [53] 2019 ↑	
	Rathnayake et al. [54] 2019 ↑	
	Miliukhina et al. [50] 2020 ↑	
	Usenko et al. [55] 2020 ↓	
Leptin	Mahlknecht et al. [45] $2012 \rightarrow$	
	Rahnemayan et al. [98] 2021 \rightarrow	
NGAL	Eidson et al. [56] 2017 ↑	
NLRP3	Chatterjee et al. [43] 2020 ↑	
	Fan et al. [44] 2020 ↑	
	Roy et al. [99] 2021 ↑	
PDGF-BB	Mahlknecht et al. [45] 2012 \uparrow	
Prolactin	Mahlknecht et al. [45] 2012 ↑	
S100A8/9 (Calprotectin)	Dumitrescu et al. [100] 2021 \uparrow	
S100B	Dumineseu et al. [100] 2021	Sathe et al. [101] 2012 ↑
SAA (Serum amyloid A)		Hall et al. [63] 2018 \uparrow
SCF	Brockmann et al. [38] 2016 ↑	Santaella et al. [65] $2020 \rightarrow$
TGF-β1	Dioekinani et al. [50] 2010	Chen et al. [60] 2018 \uparrow
TGF-α		Santaella et al. [65] 2020 \rightarrow
TIMP-2	Mahlknecht et al. [45] $2012 \rightarrow$	Santaena et al. $[03]$ 2020 \longrightarrow
		Delegado Alverado et al. [49] 2017 A
TNF-α	Koziorowski et al. [35] 2012 \uparrow	Delgado-Alvarado et al. [48] 2017 ↑
	Hu et al. [36] $2015 \uparrow$	Chen et al. [60] 2018 \longrightarrow
	Gupta et al. [51] 2016 \downarrow	Schröder et al. [46] 2018 ↑
	Wang et al. [26] 2016 ↑	Hu et al. [36, 64] 2015, 2019 \longrightarrow
	Qin et al. [25] 2016 ↑	Iwaoka et al. [61] 2020 ↑
	Csencsits-Smith et al. [58] 2016 ↑	
	Eidson et al. [56] 2017 ↑	
	Karpenko et al. [39] $2018 \rightarrow$	
	Kim et al. [27] $2018 \rightarrow$	
	Kouchaki et al. [59] 2018 ↑	
	Rocha et al. [40] 2018 ↓	
	Schröder et al. [46] 2018 \longrightarrow	
	Alrafiah et al. [41] 2019 \longrightarrow	
	King et al. [28, 29] 2019 ↑	
	Lin et al. [53] 2019 ↑	
	Rathnayake et al. [54] 2019 ↑	
	Martin-Ruiz et al. [57] 2020 ↑	
	Miliukhina et al. [50] 2020 \downarrow	
sTNFR1 and sTNFR2	Rocha et al. [83] 2014 ↑	
sVCAM1	Perner et al. [80] 2019 ↑	
YKL-40		Hall et al. [63] 2018 ↓

Table 1

 \uparrow : Significantly elevated levels in comparison to controls. \longrightarrow : No significant differences in comparison to controls. \downarrow : Significantly reduced levels in comparison to controls.

to healthy controls [36, 46, 48, 60, 61, 63–65]. A detailed overview of all assessed CSF markers along with the respective findings and references is given in Table 1.

Genetic-associated PD

LRRK2 (Leucine-rich repeat kinase 2)

There are 3 studies available which explored inflammatory markers in PD patients with *LRRK2*

mutations as well as in non-manifesting *LRRK2* mutation carriers.

Two studies with focus on *LRRK2* used serum samples from the Michael J. Fox Foundation LRRK2 Cohort Consortium and assessed 32 and 23 inflammatory markers by multiplex assays. Both studies found similar serum levels of inflammatory markers between PD patients with *LRRK2* mutations compared to those with *LRRK2*-wildtype status [38, 66]. In a subgroup with CSF available, higher levels of VEGF and IL-8 were detected in PD patients with *LRRK2* mutations compared to those with *LRRK2*-wildtype status [66]. A recent study from Israel confirmed the findings of relatively similar inflammatory profiles between PD patients with vs. without *LRRK2* mutations as the colleagues also did not find any differences in blood or CSF levels of cytokines [67].

Within the group of PD patients with *LRRK2* mutations, higher serum levels of IL-8, MCP-1 and MIP1- β seem associated with a more severe phenotype comprising cognitive impairment, orthostatic hypotension and REM-Sleep-behaviour disorder [14].

While 1 of the 3 studies reports higher serum levels of IL-1B [66], the other 2 studies did not find differences in inflammatory profiles in blood and/or CSF between non-manifesting LRRK2 mutation carriers and healthy controls, even when clinically stratified for prodromal markers [38, 67]. A recent study in Norwegian individuals revealed higher CSF levels of TNF- α in *LRRK2* mutation carriers when compared to controls [68]. Given these somewhat variable findings, one has to keep in mind that different prodromal symptoms vary in their predictive value to PD conversion and that the rate of progression from prodromal to manifest PD is age-dependent and varies among patients. Therefore, these cross-sectional results in the prodromal cohorts need to be interpreted with caution and warrant further longitudinal studies, ideally in individuals who convert to PD during study.

GBA (Glucocerebrosidase)

While one small study reports increased plasma levels of IFN- γ , IL-1 β , IL-2, and TNF- α in 8 PD patients carrying heterozygous mutations in the *GBA* gene [50], two larger studies did not find any differences in blood and CSF levels between PD patients as well as healthy individuals with vs. without heterozygous *GBA* mutations [67, 69].

PRKN/PINK1 (Parkin/PINK1)

It has been shown that dysfunction in Parkin and PINK1 enhances mitochondrial stress with activation of inflammatory processes and neurodegeneration via the STING protein cascade (stimulator of interferon genes) [70]. Consequently, PD patients with bi-allelic and heterozygous mutations in *PRKN/PINK1* showed increased serum levels of IL-6 in a gene-dosage manner in 3 cohorts [71].

We conclude that the role of inflammation in PD is reflected in increased inflammatory markers in blood and CSF in sporadic PD patients compared to healthy control individuals. Studies in genetically associated PD (*LRRK2*, *GBA*, *PRKN/PINK1*) are rare and indicate similar profiles as seen in sporadic PD. Data in at-risk and prodromal mutation carriers are sparse and conflicting and need further validation in prospectively followed longitudinal cohorts.

IS THERE A ROLE OF ANTI-INFLAMMATORY MEDIATORS IN BLOOD/CSF?

While many studies focus on pro-inflammatory profiles, it is more and more established that inflammation is a balance between pro- and antiinflammatory processes.

The best investigated anti-inflammatory marker in the context of PD is IL-10 in blood. However, reports are quite heterogenous. While some studies show higher levels in sporadic PD patients, others find no differences or lower levels in sporadic PD patients compared to healthy controls [25, 27–29, 35, 38–40, 46, 54, 57]. Two studies have been done in CSF and found no differences between sporadic PD patients and healthy controls [46, 64].

Within the group of PD patients, single studies report higher blood levels of IL-10 to be associated with motor and cognitive impairment as well as with higher CSF levels of α -synuclein [55, 57]. Whether such findings represent compensatory upregulation to keep the balance between pro-and anti-inflammatory species remains to be elucidated. A detailed overview of the respective findings in blood and CSF along with references is given in Table 1.

DO INFLAMMATORY PROFILES IN BLOOD REFLECT THOSE IN CSF INDICATIVE OF A CROSS-TALK BETWEEN PERIPHERY AND BRAIN?

When designing clinical trials, one has to decide which biomaterial should to be repeatedly collected for outcome measurements. In this context, blood is more easily accessible than CSF. However, we should know whether blood profiles are representative for profiles in CSF and can serve as proxy for centralnervous system neuroinflammation?

So far, there are 3 comprehensive studies published in sporadic PD that assessed a variety of inflammatory markers by multiplex assay in CSF/serum pairs [16, 56, 72]. The 2 smaller cohorts comprised 12 and 22 PD patients whereas our own cohort consisted of 453 sporadic PD patients. Importantly, the platform/panel (Mesoscale: V-PLEX: IFN-γ, IL-1β, IL-2, IL-4, IL-6, IL-8, IL-10, IL-12p70, IL-13, TNF-α) that was used in the 2 smaller studies was identical and fully overlapped with the inflammatory markers assessed in our own study (Myriad: AFP, BDNF, CA-125, CEA, CKMB, ENA-78, FactorVII, FABP, GH, IgE, ICAM-1, IL-1α, IL-1β IL-2, IL-3, IL-4, IL-5, IL-6, IL-7, IL-8, IL-10, IL-12p40, IL-12p70, IL-13, IL-15, IL-16, IL-18, Leptin, Lymphotactin, MDC, MIP-1β, MMP3, MMP9, MCP-1, PSA-f, SCF, TPO, TSH, TF, TNF- α , TNF- β). Results from all 3 studies are quite congruent: (I) Only a fraction of markers is reliably detectable in both CSF and serum. (II) Correlations of cytokines between blood and CSF are sparse. In the 2 smaller studies, none of the markers IL-6, IL-8, IL-10, IFN- γ , TNF- α showed a significant correlation between serum and CSF. The first study with 12 patients additionally assessed CRP which showed a good correlation between serum and CSF. In our larger study of 41 markers, ICAM-1, IL-4, IL-6, IL-12p40, MCP-1, MIP-1β, MMP3, Leptin, and TSH were correlated between CSF and serum in females and in males whereas IL-8, IL-13 and CKMB were correlated only in females.

These findings indicate that inflammatory markers in blood and CSF do not reflect one another and that they might be regulated independently of each other. This poses a challenge for clinical trials. Which biomaterial should to be collected for repeated outcome measurements – the more easily accessible blood or CSF by lumbar puncture?

DO BLOOD/CSF INFLAMMATORY PROFILES CHANGE OVER THE DISEASE COURSE AS ASSESSED IN REPEATEDLY TAKEN BIOSAMPLES?

Inflammatory profiles do not follow a linear pattern as recent evidence highlights that inflammation in PD is maximized in the early disease stages and maintains a chronic profile during the course of the disease. Knowledge of the trajectories of the different biofluid markers in relation to disease progression is important when interpreting therapeutic effects (beneficial/risk) in clinical trials targeting the immune system.

One study used a multiplex assay assessing 37 cytokines in serum of 25 sporadic PD patients every 6 months over 2 years. While levels of IL-4 increased, plasma levels of IL-2, IL-9 and IL-15 decreased over time. Other cytokines did not change significantly

over time [58]. Another study reports stable blood levels of CRP over a period of 3 years in 313 PD patients with a mean disease duration of 7.9 years at baseline. Notably, the patients have been stratified by their individual CRP levels at baseline (low, mid, high) [73]. Another study in 47 PD patients showed an increase in serum levels of C3, C4, and IL-6 in 30–50% of patients over a period of 2 years [74].

So far, 1 study reports longitudinal measurements of YKL-40 in repeatedly taken CSF samples of 63 sporadic PD patients. The authors report an increase of CSF levels of YKL-40 over 2 years in the whole PD cohort as well as in subgroups when stratified by disease duration \leq or>5 years disease duration [75].

We conclude that large longitudinal studies with repeatedly collected blood and CSF biosamples in *de-novo* patients over several years are missing. However, such data are of utmost importance and will provide information on how inflammatory profiles evolve over time and in concert with other neurodegenerative/disease-specific biomarkers.

ARE BLOOD/CSF INFLAMMATORY PROFILES ASSOCIATED WITH PHENOTYPICAL TRAJECTORIES IN PD?

In order to enrich cohorts for maximized therapeutic benefit, define clinically reasonable outcome measures and target specific clinical milestones, knowledge on the predictive/prognostic value of inflammatory profiles in relation to clinical trajectories is important.

There are several studies in blood and CSF that investigated inflammatory profiles in relation to phenotypical characteristics of motor and non-motor symptoms, the latter with focus on cognition, depression and sleep. A detailed overview of associations between inflammatory profiles in blood and CSF with phenotypical characteristics is given in Table 2 along with all references.

Inflammatory markers in blood: Phenotypical trajectories

It was repeatedly shown cross-sectionally that higher blood levels of IL-6 [42, 48, 76] and CRP [30, 77] are associated with worse motor function. Other single studies further found that higher blood levels of fractalkine [78], IL-1 β [44], IL-15 [72], IFN- γ [56], S100B [79], TNF- α [72], and VCAM1 [80] are related to worse motor function. One longitudinal study could further support the finding that

			В	lood						CSF		
	Mortality	Motor function	Cognitive function	Depression	Hallucination	Fatigue Sleep RBD	Mortality	Motor function	Cognitive function	Hallucination	Depression	Fatigue Sleep RBD
C3			Vesely [74, 85] 2018, 2019									
C4			Vesely [74] 2018									
CCL2 MCP-1								Santaella	Lerche		Lindqvist	Lindqvist [23]
CRP	Sawada [73] 2015	Umemura [81] 2015	Mollenhauer [84] 2019		Sawada [24] 2014			[88] 2020 Hall [63] 2018	[72] 2022 Lindqvist [23] 2013		[23] 2013 Lindqvist [23] 2013	2013 Lindqvist [23] 2013
		Santos-García [30] 2019 Li	Martin-Ruiz [57] 2020					Moghaddam [87] 2018	Hall [63] 2018 Moghaddam		Hall [63] 2018 Lindqvist	
CV2CL1 Encetalle		[77] 2021						Hatcher-Martir	[87] 2018		[23] 2013	
CX3CL1 Fractalki	ne	Gupta [78] 2021						[89] 2021	l			
CXCL10 IP-10												Lindqvist [23] 2013
FABP			Lerche [72] 2022						Lerche [72] 2022			
IL-1ß		Fan [44] 2020										Hu [36] 2015
IL-1RA						Herlofson [10 2018	2]					
IL-2		King [28] 2019										
IL-4		King [28] 2019										
IL-6	Dufek [86] 2015	Delgado- Alvarado [48] 2017 Green [76] 2019 Lian	Vesely [85] 2019 Martin-Ruiz [57] 2020 Usenko [55] 2020	Vesely [74] 2018		Pereira [103] 2016		Lian [42] 2019	Lindqvist [23] 2013 Yu [62] 2014			

Table 2

IL-8					Lerche	Lerche	
IL-10		Usenko [55] 2020 Martin-Ruiz	Karpenko [104] 2018		[72] 2022	[72] 2022	
IL-12p40	Yilmaz [82] 2018	[57] 2020 Yilmaz [82] 2018					
IL-15	Lerche [72] 2022						
IL-17A		Green [76] 2019	Green [76] 2019				
IL-18					Lerche [72] 2022		
IFN-y	Eidson [56] 2017	Martin-Ruiz [57] 2020					
MMP10					Santaella [88] 2020		
S100B	Carvalho [79] 2015			Carvalho [79] 2015	¥¥ 11	TT 11	¥¥ 11
SAA Serum amyloid A SCF					Hall [63] 2018	Hall [63] 2018 Lerche [72] 2022	Hall [63] 2018
TNF-α	Lerche [72] 2022	Usenko [55] 2020 Martin-Ruiz [57] 2020 Lerche				[72] 2022	
sTNFR1 sTNFR2		[72] 2022 Rocha [83] 2013					
sVCAM1	Perner [80] 2019			Herlofson [102] 2018			
YKL-40						Wennström [90] 2015 Hall [75] 2016	

Straight = positive associations meaning higher levels of the respective inflammatory marker are associated with worse clinical outcome measure. *Italic* = *Results show an inverse association meaning lower levels of the respective (anti)-inflammatory marker are associated with worse clinical outcome measure.* **Bold** = **Studies with longitudinal clinical data.** Please note that due to space limitation we only name the first author with year of publication and do not refer to co-authors with "et al.".

higher CRP levels are associated with more severe motor impairment [81]. Contrary, lower blood levels of the anti-inflammatory makers IL-4 [28] and IL-12p40 [82] were also associated with worse motor function.

Several cross-sectional studies report higher levels of CRP, FABP, IL-6, IL-10, IL-17A, TNF- α , and TNFR to be related with worse cognitive function and dementia [55, 72, 76, 83]. One longitudinal study supports the finding that higher levels of CRP are associated with cognitive decline over time [84]. Similarly, it was shown longitudinally that higher levels of C3 and C4 are related to reduced memory function [74, 85]. Another longitudinal study created an inflammatory composite score based on Patients' individual blood levels of CRP, IFN- γ , IL-6, IL-10, and TNF- α . While a higher composite score predicted worse MoCA scores over 3 years, no association was found with phenoconversion to manifest dementia [57].

Two longitudinal studies report an association between higher levels of CRP and IL-6 with overall mortality in PD [73, 86].

Inflammatory markers in CSF: Phenotypical trajectories

Two studies report higher CSF levels of CRP to be associated with worse motor function [63, 87]. Other single studies found that higher CSF levels of IL-6 [42], IL-8 [72], and SAA [63] are related to worse motor function. One longitudinal study reports that higher CSF levels of MCP-1 and MMP-10 are associated with more severe motor impairment [88]. Contrary, lower levels of the anti-inflammatory marker fractalkine and IL-18 were associated with worse motor function [72, 89].

Several cross-sectional studies report higher CSF levels of CRP and IL-6 to be related with worse cognitive function [23, 63, 87]. Other single studies further found that higher CSF levels of MCP-1, FABP, IL-8, SAA, SCF, and YKL-40 are associated with cognitive impairment [63, 72, 90]. One longitudinal study supports that higher CSF levels of YKL-40 are associated with cognitive decline over time [75].

Higher CSF levels of CRP and MCP-1 were associated with depression and fatigue [23, 63].

We conclude that higher blood and CSF levels of inflammatory markers are associated with more severe clinical trajectories of motor and non-motor symptoms cross-sectionally as well as longitudinally (Fig. 1).

ARE BLOOD/CSF INFLAMMATORY PROFILES ASSOCIATED WITH CSF LEVELS OF NEURODEGENERATIVE/ PD-SPECIFIC BIOMARKERS?

In order to map the complex picture of neurodegeneration with its different pathway-related endpoints (α -synuclein aggregation, concomitant amyloid-beta (A β) and tau pathology) and (neuro)inflammation, comprehensive biomarker analyses are needed.

The 3 studies that assessed inflammatory markers by multiplex assay in CSF/serum pairs (described above) also performed additional analyses in relation to CSF levels of neurodegenerative/PD-biomarkers [16, 56, 72].

Inflammatory markers in blood: CSF levels of neurodegenerative/PD-specific biomarkers

In our own study, we could show that higher blood levels of ICAM-1, IL-18, and MIP-1 β are associated with higher CSF levels of t-Tau and p181-Tau. Higher blood levels of FABP, MMP3, TF, and TNF- α are correlated with higher CSF levels of NFL. Higher blood levels of IL-10, IL-12p40, IL-13, IL-16, IL-18, MCP-1, and MIP-1 β are associated with higher CSF levels of α -synuclein. Notably, the majority of these associations were only found in female PD patients [72].

Inflammatory markers in CSF: CSF levels of neurodegenerative/PD-specific biomarkers

In the study by Wijeyekoon, higher CSF levels of IL-1 β and IL-8 were associated with higher CSF levels of t-Tau whereas no correlation was seen between CSF inflammatory markers with p-Tau, α -synuclein or A β_{1-42} [16]. Our own study revealed that higher CSF levels of FABP, ICAM-1, MMP3, SCF, and TF were associated with higher CSF levels of t-Tau, p181-Tau and α -synuclein in both, males and in females whereas the same association was detected with A β_{1-42} only on males [72].

There is only 1 longitudinal study with repeatedly taken CSF samples over 2 years available. Here, higher baseline CSF levels of YKL-40 were associated with higher baseline CSF levels of t-Tau, p-Tau and α -synuclein but not with A β_{1-42} . Similarly, an increase in CSF levels of YKL-40 over 2 years was associated with an increase in CSF levels of t-Tau, p-Tau, and α -synuclein [75]. In conclusion, the studies indicate that associations between inflammatory markers with CSF levels of neurodegenerative/PD-biomarkers are primarily seen with t-Tau, p-Tau, and α -synuclein but not with A β_{1-42} .

CONCLUSION

Based on findings of the studies discussed in this review we conclude: (I) The role of inflammation in PD pathogenesis is reflected in increased inflammatory markers in blood and CSF in sporadic PD patients compared to healthy control individuals. (II) Within the group of PD patients, higher blood and CSF levels of inflammatory markers are associated with more severe clinical trajectories of motor and non-motor symptoms cross-sectionally as well as longitudinally. (III) In line with evidence from postmortem and cell studies, the kind of blood and CSF levels of inflammatory markers indicate a contribution of the innate and adaptive immune system in both CNS and periphery. (IV) Correlations of inflammatory markers in blood and CSF are sparse indicating that they do not reflect one another. This poses a challenge for clinical trials as one has to decide which biomaterial should to be collected for repeated outcome measurements-the more easily accessible blood or CSF by lumbar puncture.

Inflammation is also relevant in the pathogenic mechanisms in other neurodegenerative diseases, among them also Alzheimer's disease (AD). Interestingly, similar metabolites as reported relevant in PD patients are also elevated in blood and/or CSF in AD patients when compared to healthy controls: CRP, IL-16, IL-2, IL-4, IL-6, IL-10, IL-12, IL-18, MCP-1, MCP-3, IL-8, IFN-γ, TGF-β, TNA-α, and YKL-40 [60, 91–95]. Thus, inflammatory processes seem to be somewhat disease-unspecific and triggered by different disease-related protein species such as α synuclein, $A\beta$, and tau. These findings in turn offer the chance to accumulate knowledge on inflammation across different neurodegenerative diseases. We acknowledge the following shortcomings: (I) Longitudinal studies with repeatedly collected blood and CSF biosamples in de-novo patients over several years, ideally accompanied by other disease-related biomarkers are missing. This will provide information on how inflammatory profiles evolve over time and in concert with other neurodegenerative/diseasespecific protein levels and on the predictive value for the development of disease-related milestones such

as dementia. (II) Studies in (genetically) at-risk and prodromal individuals are just beginning to emerge. So far, no robust conclusions can be drawn as longitudinal data in phenoconverters are missing. (III) The type of assays used (single vs. multiplex, platforms), of cytokines assessed and the cohorts' characteristics (samples size, disease duration) are quite variable. This limits the informative value of each individual marker and comparability across studies. (IV) There is not the one single inflammatory marker in blood and CSF that is associated with clinical trajectories, disease progression and neurodegenerative/diseasespecific protein levels. The most frequently reported markers in this context are IL-1B, IL-6, IL-8, MCP-1, and TNF- α (Fig. 1). However, these are also the most investigated ones so that there might be a bias in interpretation. An elegant way might be a composite score out of the 5-10 top markers and thereby stratifying patients by their intra-individual inflammatory load.

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CONFLICT OF INTEREST

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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