Current Journal of Neurology

Original Paper



Curr J Neurol 2024; 23(1): 21-38

New diagnosis of multiple sclerosis in the setting of recent Sinopharm COVID-19 vaccine (BBIBP-CorV) exposure: A series of clinical cases and updated review of the literature

Received: 05 Sep. 2023 Accepted: 06 Nov. 2023

Sepideh Paybast¹, Melika Jameie¹, Mojtaba Shahbazi¹, Mohammad Amin Habibi², Seyed Ehsan Mohammadianinejad¹, Mohammad Hossein Harirchian¹

¹ Iranian Center of Neurological Research, Neuroscience Institute, Imam Khomeini Hospital Complex, Tehran University of Medical Sciences, Tehran, Iran

² Student Research Committee, School of Medicine, Qom University of Medical Sciences, Qom, Iran

Keywords

COVID-19; COVID-19 Vaccines; Multiple Sclerosis; CNS Demyelinating Autoimmune Diseases; Safety; Demyelinating Diseases

Abstract

Background: Multiple sclerosis (MS) is the most common cause of non-traumatic disability in young individuals. There are limited reports of developing demyelinating events following the coronavirus disease 2019 (COVID-19) vaccination.

Methods: We reported all individuals (n = 8) with new MS diagnoses with recent exposure (\leq 6 weeks) to the Sinopharm (BBIBP-CorV) vaccine between September 2021 and June 2022. We also reviewed the related literature published as of September 2023.

Results: Of 338 newly diagnosed patients with MS who

attended our tertiary referral MS center during the study period, 8 (2.36%) had their first demyelinating attack with a median interval of 2 [2.0, 4.0] weeks following the Sinopharm vaccine (sex ratio 1:1, median age: 20.5 [18.0, 27.0] years). No personal or family history of autoimmune/neurological disorders was documented, except for one patient's history of a previous potential demyelinating event and another's family history of immune thrombocytopenic purpura (ITP).

How to cite this article: Paybast S, Jameie M, Shahbazi M, Habibi MA, Mohammadianinejad SE, Harirchian MH. New diagnosis of multiple sclerosis in the setting of recent Sinopharm COVID-19 vaccine (BBIBP-CorV) exposure: A series of clinical cases and updated review of the literature. Curr J Neurol 2024; 23(1): 21-38.

Copyright © 2024 Iranian Neurological Association, and Tehran University of Medical Sciences Published by Tehran University of Medical Sciences

Corresponding Author 1: Mohammad Hossein Harirchian Email: harirchm@tums.ac.ir

Corresponding Author 2: Seyed Ehsan Mohammadianinejad Email: ehsanneuro@gmail.com

This work is licensed under a Creative Commons Attribution-NonCommercial 4.0 international license (http://creativecommons.org/licenses/by-nc/4.0/). Non-commercial purposes uses of the work are permitted, provided the original work is properly cited.

All patients had demyelinating brain MRI lesions, and 4 had cervical spinal cord involvement. The brain areas most commonly affected were the periventricular and subcortical regions. Positive oligoclonal bands (OCBs) in all patients supported the MS diagnosis. All patients were diagnosed with relapsing-remitting MS and received intravenous methylprednisolone (IVMP) alone or in combination with plasma exchange (3/8). Rituximab was the most frequently used diseasemodifying treatment (3/8).

Conclusion: This study provides preliminary evidence of a potential association between the Sinopharm vaccine and the initial manifestations of MS. However, further larger-scale studies with control groups and long-term follow-ups are needed to confirm this association and determine the underlying mechanisms.

Introduction

(COVID-19), Coronavirus disease 2019 а highly contagious disease caused by the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), was initially reported in December 2019.1 Since the onset of the COVID-19 pandemic,² it has been linked to significant mortality and morbidity,³⁻⁷ profoundly impacting the mental and physical well-being of both patients and healthcare professionals.⁸⁻¹⁰ There have also been reports of associated neurological manifestations.¹¹ In response to these challenges, vaccination has emerged as the most effective strategy for the mitigation of the associated mortality and morbidity.12

Multiple sclerosis (MS) is a demyelinating inflammatory disease of the central nervous Genetic system (CNS).13 variants and environmental factors, including vitamin D deficiency, obesity, smoking, and the Epstein-Barr virus are the factors associated with MS.14,15 Notably, several studies have explored the potential involvement of viral infections in the pathogenesis of MS.15-18 Recent reports on the co-occurrence of SARS-CoV-2 infection and neuroimmunological disorders have raised questions about their potential association.¹⁹⁻²¹ A similar discourse has emerged concerning the of SARS-CoV-2 vaccination impact on neuroimmunological disorders.22-30

The relationship between vaccines and the potential risk of demyelinating diseases has long been an area of research interest,³¹⁻³⁶ with controversy surrounding this association.^{34,35} However, except for the yellow fever vaccine,³⁶ there is no sufficient evidence to indicate a causal

association between most vaccines, including hepatitis B, human papillomavirus (HPV), influenza, and "measles, mumps, and rubella" (MMR), and MS activity.^{31,33-35} Recently, concerns regarding neurological adverse events following immunization (AEFI) with SARS-CoV-2 vaccines have arisen,^{26-28,37,38} including limited reports of the CNS demyelinating events potentially associated with these vaccines.^{22-25,28,30,39-43}

In this case series, we have reported 8 individuals who experienced the initial manifestations of MS within a temporal relationship (≤ 6 weeks) with the Sinopharm vaccine exposure. This case series is unique for two primary reasons. Firstly, while studies evaluating MS relapse following SARS-CoV-2 vaccination among patients with established MS have been conducted,^{29,44,45} reports regarding the new onset of the disease in seemingly healthy individuals still remain limited. Secondly, the existing literature predominantly concentrates on vaccines other than the Sinopharm (BBIBP-CorV) vaccine.22-25,28,30,39-41 Notably, except for one case series,⁴⁶ we are not aware of any previously published article concerning a new diagnosis of MS after vaccination with Sinopharm (BBIBP-CorV).

Materials and Methods

Study design and ethics statement: This is a single-center case series of prospectively collected data on 8 consecutive individuals with their first CNS demyelinating event following SARS-CoV-2 vaccination. These individuals attended an academic hospital affiliated with the Tehran University of Medical Sciences, Tehran, Iran, between September 2021 and June 2022. This study was approved by the institutional review board (IRB) of the Tehran University of Medical Sciences, Tehran, Iran, and followed the CAse REports (CARE) guideline.⁴⁷ According to the declaration patients' of Helsinki, anonymity and confidentiality were protected, and informed consent was obtained.48

Study population: All individuals attending the hospital (whether they were referred to the hospital or self-referred) during the study period with the following criteria were eligible for inclusion: (a) new onset CNS demyelination symptoms, (b) SARS-CoV-2 vaccination in the past 6 weeks, lack of infection with SARS-CoV-2 during this interval, (c) a neurologist-confirmed diagnosis of MS.⁴⁹ A 6-week time frame was chosen based on the typical time frame suggested

for neurological AEFI (Table 1).⁵⁰ No limitations were imposed regarding patients' age, or the type of vaccine received.

Study measures: The following information was collected by a neurology resident (M.SH.) and double-checked by 3 MS specialists (S.P., MH.H., and SE.M.) to validate the accuracy of the collected data: (a) demographics, past medical history (PMH), familial, and habitual history through the in-person interview, (b) vaccinerelated characteristics according to the COVID-19 vaccination cards (type, dose, and the received date), (c) MS-related clinical, laboratory, and neuroimaging characteristics **[initial** presentations and severity of MS attack, the interval between vaccination and the disease onset, cerebrospinal fluid (CSF) oligoclonal bands (OCBs) and IgG index, as well as brain, cervical spine, and thoracic spine magnetic resonance imaging (MRI) findings], (d) acute phase management, and the chosen disease-modifying therapy (DMT).

MS diagnosis was established based on the McDonald criteria 2017 and confirmed by an expert in the field, according to the presence of typical MRI lesions and positive OCBs, as well as the absence of clinical or MRI red flags supporting alternative diagnoses.⁴⁹ Attack severity was defined based on increase in the Expanded Disability Status Scale (EDSS) on the day of maximal worsening: mild (< 1.0), moderate (1.0-2.5), and severe (\geq 3.0).⁵¹

Quantitative and qualitative data are presented as median (interquartile range [IQR]), and number (percentage), respectively.

Results

Of the 338 newly diagnosed patients with MS during the study period, 8 (2.36%) had their first MS demyelinating attack in temporal association with the Sinopharm vaccine. Moreover, 4 of the 8 patients were women, and their median age was 20.5 [18.0, 27.0] years. Except for a history of a probable previous demyelinating event in 1 patient and а familial history of immune thrombocytopenic purpura (ITP) in another, no personal or familial history of autoimmune or neurological conditions was reported. In addition, 6 patients reported symptom onset after the 2nd vaccine dose, and the median interval between the $1^{st}(v1)/2^{nd}(v2)$ dose and symptom initiation was 2.0 [2.0, 4.0] weeks. Initial presentations varied from mild sensory syndrome to severe multifocal disseminated demyelination. Attack severity was moderate in most patients (5/8).

Brain MRI revealed demyelinating lesions in all (8/8) patients. Periventricular and subcortical regions were the most frequently affected brain regions, with lesions observed in 7 and 6 patients, respectively. Other affected brain regions include juxtacortical (3/8), cortical (2/8), and the brain stem (3/8). Spinal cord involvement was seen in 4, with the upper and lower cervical spine involvement in 2 and 2 patients, respectively. Gadolinium-enhanced lesions were observed in 4 patients (patients 2, 5, 6, and 8). Notably, apart from 1 patient (patient 3) who exhibited multiple hypointense lesions in T1 sequences (black holes), no other patients displayed such findings in their T1 images.

Table	1.	Suggested	criteria	for	labelling	causality	in	immune-associated	neurological	adverse	events	following
immun	izat	tion (AEFI)	suggeste	ed by	y Butler et	al. ⁵⁰						

Causality	Time fra	ame [*]	Risk factor		Alternative etiology	Notes
Probable	< 6 week	AND	No risk factors	AND	No indication of an	Ruling out other risk
					alternative etiology	factors or etiologies
						identified by clinical,
						laboratory, radiological
						and electrophysiological
						assessment, as indicated
Possible	6-12	AND/	There may be an	AND	There may be an	e.g., presence of a
	week	OR	indication of an	/OR	indication of an	previous episode of
			alternative etiology,		alternative risk	Bell's palsy in a patient
			but unlikely to		factor, but unlikely to	with post-vaccination
			explain the event		explain the event	Bell's palsy
Unlikely	< 24 hour	AND/	Alternative etiology	AND	Alternative risk	e.g., Campylobacter
	or > 12	OR	fully explains	/OR	factor fully explains	diarrhea preceding
	week		the event		the event	Guillain-Barré syndrome

AEFI: Adverse events following immunization

*Time interval between immune-associated AEFI and vaccination

Thoracic MRIs were unremarkable in all patients. Positive OCBs supported MS diagnosis in all patients. With a relapsing-remitting MS (RRMS) diagnosis, all patients received intravenous methylprednisolone (IVMP) with oral tapering. Of the 8 patients, 3 required therapeutic plasma exchange (TPE). All patients achieved marked clinical recovery. Rituximab (3/8) and dimethyl fumarate (DMF) (2/8) were the most frequently prescribed DMTs (Table 2).

According to the criteria proposed by Butler et al. for labeling causality in immune-associated neurological AEFI⁵⁰ (Table 1), 7 out of 8 patients were classified as "probably" related (AEFI occurring within 6 weeks after vaccination with no indication of an alternative etiology or no risk factors). Patient number 3 was deemed "unlikely" to be associated with the vaccine, since, although the event occurred within 6 weeks, it could be explained by an alternative etiology and/or risk factors.

Patient 1 was a 17-year-old man with a 5-day history of left facial numbness, initiated 2 weeks after v2. Neurological examination revealed decreased light touch sensation in the facial area innervated by the left trigeminal nerve's second (V2) and third (V3) divisions, and an EDSS score of 1.0. CSF analysis revealed 4 OCBs. Serum and CSF evaluations were negative for inflammatory, metabolic, and infectious diseases. Neuromyelitis optica-antibody (NMO-antibody) and anti-myelin oligodendrocyte glycoprotein (anti-MOG) IgG were negative. Brain MRI demonstrated a few nonenhancing ovoid T2 periventricular and subcortical lesions and 1 cortical lesion.

A cervical MRI revealed no specific findings. With RRMS diagnosis, IV-MP (1 gr/day, 3 days) was given, leading to a complete symptom recovery. DMF was then initiated (120 mg daily, titrated to 240 mg twice a day).

Patient 3 was a 39-year-old female smoker with a 7-day history of right upper limb paresis 2 weeks after v2. She reported a history of 3 days of right facial numbness 15 years ago. Neurological examination revealed generalized hyperreflexia and right upper limb mono-paresis (power 3/5 according to the Medical Research Council [MRC] muscle strength scale). EDSS was estimated to be 2.5. Extensive laboratory assessments revealed no abnormal findings in the serum and CSF except for 5 OCBs restricted to the CSF. The brain and cervical spine MRIs demonstrated slight, generalized atrophy, multiple non-enhancing hyperintense lesions, predominantly in juxtacortical and periventricular locations, and a short-segment, non-enhancing C2 lesion (Figure 1). Moreover, multiple black hole lesions were observed in T1 sequences. With a diagnosis of RRMS, IV-MP (1 gr/day, 5 days) was commenced, followed by rituximab (1 gr biannually). A previous history of facial numbness and MRI alterations (atrophy and black holes) suggest a latent demyelinating process initiated years ago. She achieved a significant recovery and was discharged with an EDSS score of 1.0.

Patient 4 was a 29-year-old male smoker with left leg debility initiated a week after v2. Neurological examination exhibited a muscle strength of 4/5 in his left lower limb with an upward plantar reflex and EDSS score of 2.5. CSF analysis demonstrated a mildly elevated IgG index (0.78) and 9 unique OCBs. Brain MRI revealed a few non-enhancing periventricular and subcortical demyelinating lesions. A cervical MRI revealed 2 short-segment lesions without gadolinium enhancement. With a diagnosis of RRMS, IV-MP (1 gr/d, 3 days) was started, markedly improving his symptoms. Subsequently, fingolimod (0.5 mg) was started under 6 hours of cardiac monitoring.

Patient 5 was a 14-year-old girl with acute diplopia 2 weeks after v1. Neurological examination revealed right 6th nerve palsy, with an EDSS of 1.5. Except for 11 unique CSF OCBs, serum and CSF analysis did not indicate other abnormalities. Brain MRI showed numerous ovoid T2 periventricular, subcortical, cortical, and brain stem lesions, some of which showed abnormal gadolinium enhancement. The cervical spine MRI was normal (Figure 2). With a diagnosis of RRMS, IV-MP (1 gr/day, 5 days) and TPE were started (5 exchanges daily with 1.5-liter plasma volume, using 5% albumin as a replacement fluid). Rituximab was chosen as DMT (1 gr biannually). She achieved a complete recovery at discharge.

Patient 6 was a 16-year-old girl with acute onset painful blurred vision 5 weeks after v2. Neurological examination revealed a reduced right-side visual acuity of "light perception" with a relative afferent pupillary defect (RAPD). Fundal examination showed pink disks with sharp margins. CSF analysis revealed an elevated IgG index (0.92) and 6 OCBs on isoelectric focusing.

Brain MRI revealed a few T2 subcortical, midbrain, and MCP lesions, one of which demonstrated nodular gadolinium enhancement.

Variable	Case 1	Case 2	Case 3	Case 4
Age	17	25	39	29
Sex	М	Μ	F	М
PMH	Neg	Neg	Facial numbness	Neg
HH	Neg	Neg	Smoking (20 PY)	Alcohol, Smoking (15 PY)
FH	Neg	Neg	Neg	Neg
Vaccine	Sinopharm	Sinopharm	Sinopharm	Sinopharm
Vaccine dose	v2	v1	v2	v2
Interval*	2 W	4 W	2 W	1 W
Manifestation	Lt. facial numbness	Quadri-paresthesia	Rt. upper limb paresis	Lt. lower limb paresis
Attack severity	Mild	Moderate	Moderate	Moderate
Laboratory	4 OCBs (CSF)	6 OCBs (CSF)	5 OCBs (CSF)	Elevated IgG index, 9 OCBs (CSF)
Brain MRI	Few non-enhancing PV	One enhancing lesion in	Slight generalized atrophy, Multiple	Few non-enhancing PV and SC
	and SC lesions, One	the Rt. MCP, Few non-	non-enhancing lesions, predominantly in	lesions
	cortical lesion	enhancing PV lesions	JC and PV, Multiple black hole lesions in	
			T1 sequences imaging	
Cervical MRI	Normal	One ss lower cervical,	One ss C2 non-enhancing	Two ss non-enhancing
		non-enhancing lesion	lesion	lesions
Thoracic MRI	Normal	Normal	Normal	Normal
Acute management	IVMP (3 gr)	IVMP (5 gr)	IVMP (5 gr)	IVMP (3 gr)
DMT	DMF	DMF	RTX	FG
Causality**	Probable	Probable	Unlikely	Probable

Table 2. Part I: Demographic, vaccine-related, MS-related, and management-related characteristics of patients with a new diagnosis of MS

Table 2. Part II: Demographic, vaccine-related, MS-related, and management-related characteristics of patients with a new diagnosis of MS

Variable	Case 5	Case 6	Case 7	Case 8
Age	14	19	20	21
Sex	F	F	М	F
PMH	Neg	Neg	Neg	Neg
HH	Neg	Neg	Smoking (15 PY)	Neg
FH	Neg	Neg	ITP	Neg
Vaccine	Sinopharm	Sinopharm	Sinopharm	Sinopharm
Vaccine dose	v1	v2	v2	v2
Interval [*]	2 W	5 W	2 W	4 W
Manifestation	Acute diplopia, Rt. 6 th	Acute onset blurred vision,	Acute onset blurred	Rt. facial numbness, Impaired
	nerve palsy	Rt. declined VA (light	vision, Rt. Declined VA	ocular movement, Rt. 6 th nerve
		perception), RAPD ⁺	(6/10), RAPD ⁺	palsy, Walking difficulty, Ataxia
Attack severity	Moderate	Severe	Moderate	Severe
Laboratory	11 OCBs (CSF)	IgG index ↑, 6 OCBs (CSF)	5 OCBs (CSF)	6 OCBs (CSF)
Brain MRI	Numerous PV, SC, cortical, and	Few SC, midbrain, MCP lesions,	Few non-enhancing	Two JC lesions, Numerous PV and
	brain stem lesions, Some of	One of which demonstrated	lesions within the PV, JC,	SC lesions, some of which
	which enhanced with gadolinium	nodular gadolinium enhancement	and SC	enhanced with gadolinium

MS onset following Sinopharm COVID-19 vaccination

		, 0		6
Variable	Case 5	Case 6	Case 7	Case 8
Cervical MRI	Normal	Normal	Normal	Two non-enhancing, ss
				upper cervical lesions
Thoracic MRI	Normal	Normal	Normal	Normal
Acute management	IVMP (5 gr), TPE (5 sessions)	IVMP (7 gr), TPE (5 sessions)	IV-MP (5 gr)	IV-MP (5 gr), TPE (5 sessions)
DMT	RTX	GA	INF-β 1a	RTX
Causality**	Probable	Probable	Probable	Probable
00T 0 1 1 1 7 1				a. a

Table 2. Part II: Demographic, vacc	cine-related, MS-related, ar	d management-related characteristics of	patients with a new diagnosis of MS (c	continue)
-------------------------------------	------------------------------	---	--	-----------

CSF: Cerebrospinal fluid; DMF: Dimethyl fumarate; DMT: Disease-modifying treatment; F: Female; FG: Fingolimod; FH: Familial history; GA: Glatiramer acetate; HH: Habitual history; INF- β 1a: Interferon-beta 1a; ITP: Immune thrombocytopenic purpura; IV: Intravenous; IVMP: Intravenous methylprednisolone; JC: Juxtacortical; Lt: Left; M: Male; MCP: Middle cerebellar peduncle; MP: Methylprednisolone; MRI: Magnetic resonance imaging; MS: Multiple sclerosis; Neg: Negative; OCB: Oligoclonal band; PMH: Past medical history; PV: Periventricular; PY: Pack-years; RAPD: Relative afferent pupillary defect; Rt: Right; RTX: Rituximab; SC: Subcortical; ss: Short segment; TPE: Therapeutic plasma exchange; VA: Visual acuity; v1: 1st vaccine does; v2: 2nd vaccine dose; W: weeks

*The interval between MS symptoms onset and SARS-CoV-2 vaccination

**According to the causality criteria suggested by Butler et al.⁵⁰



Figure 1. Patient 3; (a, b) multiple ovoid hyperintense lesions predominantly in juxtacortical (red arrows) and periventricular (yellow arrows) locations in axial T2-weighted fluid-attenuated inversion recovery (FLAIR) (a) sagittal T2, (b) sequences, (c) small T2 higher cervical hyper-intense lesion (green arrow) in the sagittal image



Figure 2. Patient 5; (a) multiple ovoid hyperintense lesions in juxtacortical (red arrow), subcortical (green arrows), and periventricular (yellow arrows, third and fourth ventricle) locations in axial T2-weighted fluid-attenuated inversion recovery (FLAIR), and one tumefactive-like lesion (black arrow), (b) evidence of gadolinium enhancement in some of the lesions (white arrows)

Orbital and cervical MRIs were unrevealing (Figure 3). The patient was diagnosed with RRMS, with an EDSS of 3.0, and initially treated with IV-MP (1 gr/day, 7 days), which failed to improve the visual acuity. Subsequently, she underwent 5 full-volume TPE courses, significantly improving her visual acuity (7/10). Glatiramer acetate was then initiated (40 mg/ml).

Patient 7 was a 20-year-old male smoker with acute blurred vision and pain in eye movements 2 weeks after v2. Familial history was positive for ITP. Neurological examinations revealed reduced visual acuity (6/10) and RAPD on the left side (EDSS of 1.5). Extensive evaluation covering the differential diagnostic possibilities was normal except for 5 unique CSF OCBs.



Figure 3. Patient 6; (a) few T2 midbrain (purple arrow) lesions, (b) evidence of nodular gadolinium enhancement in the tumefactive lesion (black arrow)

Brain MRI demonstrated a few demyelinating lesions within the periventricular, juxtacortical, and subcortical locations without gadolinium enhancement. Orbital and cervical MRIs were unrevealing. With RRMS diagnosis, he underwent a course of 3 g IV-MP (1 gr/day), leading to a complete clinical recovery. Interferon-beta 1a (INF- β 1a) was the chosen DMT (44 mg/ml).

Patient 8 was a 21-year-old woman with progressive right facial numbress, impaired ocular movement, and walking difficulty for 3 days. developed 4 weeks Symptoms after v2. Neurological examinations indicated right-side facial numbness, right abducens (VI) nerve palsy, generalized hyperreflexia, right ataxia, and positive left-side Babinski sign, with an EDSS of 4.0. Routine biochemistry, CSF assay, and autoimmune-related antibodies revealed no abnormalities except 6 CSF showed 2 juxtacortical OCBs. Brian MRI demyelinating lesions, as well as numerous periventricular and subcortical T2 demyelinating lesions, some of which were enhanced with gadolinium. A cervical MRI revealed two nonenhancing, short-segment upper cervical plaques. With a diagnosis of RRMS, she received IV-MP (1 gr/day, 5 days) along with 5 full-volume courses of TPE, which improved her symptoms considerably, albeit not completely (EDSS score of 1.0). DMT was planned with rituximab (1 gr biannually).

Discussion

In this series of clinical cases, we have reported 8 patients who had presented the first manifestations of MS in a close temporal relationship (≤ 6 weeks) to receiving the Sinopharm (BBIBP-CorV) vaccine.

The Sinopharm (BBIBP-CorV) vaccine, which is the most widely available SARS-CoV-2 vaccine in our country, Iran,⁵² obtained the Emergency Used Listing in May 2021.^{12,53} A retrospective cohort study, based on self-reported data from 517 vaccinated and 174 unvaccinated Iranian patients with MS, suggested that the BBIBP-CorV vaccine does not appear to affect short-term MS activity.45 Although studies have shown acceptable SARS-CoV-2 vaccine safety profiles, 53-58 concerns regarding neurological AEs have begun to emerge, with vaccine-attributed headaches being the most frequently reported.59 However, more serious neurological AEs, including Guillain-Barré syndrome,⁶⁰ Bells' palsy,⁶¹ cerebral venous sinus thrombosis, ischemic stroke, and convulsive disorders, were also reported following SARS-CoV-2 vaccination.⁶² A self-controlled case series on nearly 32 million vaccine receivers suggested an increased risk of some neurological AEFI, including Guillain-Barré syndrome, Bells' palsy, and hemorrhagic stroke, within 28 days of receiving ChAdOx1nCoV-19 or BNT162b2.²⁷ Remarkably, the author found no association between these vaccines and hospital admissions due to acute CNS demyelinating events.²⁷

A general PubMed search using the following search query ((covid 19 vaccines [MeSH Terms]) AND ((multiple sclerosis [MeSH Terms]) OR (Demyelinating Autoimmune Diseases, CNS [MeSH Terms]))) AND ((((("initial"[Title/Abstract]) OR ("first"[Title/Abstract])) OR ("new"[Title/Abstract])) OR ("onset"[Title/Abstract])) OR ("new onset"[Title/Abstract])) revealed that as of September 5, 2023, there are very few studies specifically addressing the new diagnosis of MS following SARS-CoV-2 vaccination, and are primarily limited to case reports and case series. 22-25,30,40,42,43,46 A systematic review of patients with CNS demyelinating events after SARS-CoV-2 vaccination suggested that MS-like presentations were among the most frequently reported.²⁸ According to the authors' findings up until September 2021, 12 individuals were reported with MS-like presentations following SARS-CoV-2 vaccination, of which 6 were MS relapses and 4 were diagnosed with the first MS episode without any previous history of neurological dysfunction.²⁸ In a more recent case-report-based systematic review, the authors presented 11 reported individuals with new diagnoses of MS following SARS-CoV-2 vaccination as of March 1st, 2022.63 For comprehensive insights, in table 3, we have provided detailed characteristics of 7 related studies published as of September 5, 2023, collectively encompassing a total of 26 reported cases with new MS diagnoses following immunization with SARS-CoV-2 vaccines. 22-25, 30, 40, 42, 43, 46

Most of the patients (22/26) were women and the age at MS diagnosis ranged from 19 to 66 years. PMH was reported in all patients except 3 (p5 in the study by Toljan et al.,²² p2 and p4 in the study by Nistri et al.⁴³). Accordingly, 9/26 patients had no remarkable PMH, 1/26 had unknown background,²⁴ and 7/21 had a previous history of non-neurological conditions (hypothyroidism [n = 2],^{30,46} hyperthyroidism [n = 1],⁴⁶ thyroid cancer [n = 1],⁴² hypertension [n = 1],⁴⁶ asthma [n = 1],⁴⁶ obsessive-compulsive disorder [n = 3],⁴⁶ depression [n = 1],⁴⁶ and iron-deficiency anemia [n = 3]⁴⁶).

Fujimori et al. 25 Havla et al. 24Case report1F40 40recovered Lt. facial palsy (4 years ago) FH: N/A Unknown background likely pre-existing subclinical inflammatory, CNS disease, unremarkable history of previous relapses, FH: MS in a paternal cousinBNT162b2 (v2)2w BNT162b2 (v1)Khayat-Khoei et al. 23Case series [‡] (n = 7)p2F26PMH: Neg FH: N/AmRNA-1273 (v2)14dMathew and John ⁴¹ Case report1F24PMH: Lt. facial numb., Lt. upper limb weakness (4 years ago, self-improved in a week), FH: N/ABNT162b2 (v2)5dNistri et al. 43Case seriesp2F48PMH: N/AAZD1222 (v1)8d	Study	Design	Ν	Sex	Age (year)	PMH, FH	Vaccine type (dose)	Int.
Havla et al. 24Case report1F28Unknown background likely pre-existing subclinical inflammatory, CNS disease, unremarkable history of previous relapses, FH: MS in a paternal cousinBNT162b2 (v1)6dKhayat-Khoei et al. 23Case series‡ (n = 7)p2F26PMH: Neg FH: N/AmRNA-1273 (v2)14dMathew and John ⁴¹ p5M33PMH: Neg FH: N/ABNT162b2 (v2)1dMathew Nistri et al. 43Case series p2p2F48PMH: Lt. facial numb., Lt. upper limb weakness (4 years ago, self-improved in a week), FH: N/AAZD1222 (v1)8d	Fujimori et al. ²⁵	Case report	1	F	40	recovered Lt. facial palsy (4 years ago) FH: N/A	BNT162b2 (v2)	2w
$\begin{array}{c ccccc} CNS \ disease, \ unremarkable \ history \ of \ previous \ relapses, \\ FH: \ MS \ in \ a \ paternal \ cousin \\ FH: \ MS \ in \ a \ paternal \ cousin \\ \hline FH: \ MS \ in \ a \ paternal \ cousin \\ \hline FH: \ MS \ in \ a \ paternal \ cousin \\ \hline FH: \ N/A \\ \hline \\ Mathew \\ Mathew \\ Mathew \\ Case \ report \ 1 \\ Fh \\ Nistri \ et \ al. \ A \\ \hline \\ Nistri \ et \ al. \ A \\ \hline \\ FH \\ Mathew \\ Case \ series \\ p2 \\ F \\ \hline \\ Fh \\ Fh \\ Fh \\ Fh \\ Fh \\ Fh \\$	Havla et al. ²⁴	Case report	1	F	28	Unknown background likely pre-existing subclinical inflammatory,	BNT162b2 (v1)	6d
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$						CNS disease, unremarkable history of previous relapses,		
Khayat-Khoei et al. 23Case series‡ $(n = 7)$ p2F26PMH: Neg FH: N/AmRNA-1273 (v2)14dp5M33PMH: Neg FH: N/ABNT162b2 (v2)1dMathew and John ⁴¹ Case report1F24PMH: Lt. facial numb., Lt. upper limb weakness (4 years ago, self-improved in a week), FH: N/AAZD1222 (v2)5dNistri et al. 43Case seriesp2F48PMH: N/AAZD1222 (v1)8d						FH: MS in a paternal cousin		
et al. 23 (n = 7)FH: N/ABNT162b2 (v2)1dp5M33PMH: Neg FH: N/ABNT162b2 (v2)1dMathew and John ⁴¹ Case report1F24PMH: Lt. facial numb., Lt. upper limb weakness (4 years ago, self-improved in a week), FH: N/AAZD1222 (v2)5dNistri et al. 43 Case seriesp2F48PMH: N/AAZD1222 (v1)8d	Khayat-Khoei	Case series [‡]	p2	F	26	PMH: Neg	mRNA-1273 (v2)	14d
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	et al. ²³	(n = 7)				FH: N/A		
Mathew Case report 1 F 24 PMH: Lt. facial numb., Lt. upper limb weakness (4 years ago, and John ⁴¹ AZD1222 (v2) 5d Nistri et al. ⁴³ Case series p2 F 48 PMH: N/A AZD1222 (v1) 8d			p5	Μ	33	PMH: Neg	BNT162b2 (v2)	1d
MathewCase report1F24PMH: Lt. facial numb., Lt. upper limb weakness (4 years ago, self-improved in a week), FH: N/AAZD1222 (v2)5dand John ⁴¹ 5555555Nistri et al. ⁴³ Case seriesp2F48PMH: N/AAZD1222 (v1)8d						FH: N/A		
and John ⁴¹ self-improved in a week), FH: N/ANistri et al. ⁴³ Case series p2 F48PMH: N/AAZD1222 (v1)8d	Mathew	Case report	1	F	24	PMH: Lt. facial numb., Lt. upper limb weakness (4 years ago,	AZD1222 (v2)	5d
Nistri et al. ⁴³ Case series p_2 F 48 PMH: N/A AZD1222 (v1) 8d	and John ⁴¹					self-improved in a week), FH: N/A		
	Nistri et al.43	Case series	p2	F	48	PMH: N/A	AZD1222 (v1)	8d
$(n = 16)^{##}$ FH: N/A		$(n = 16)^{\#}$				FH: N/A		
p4 F 66 PMH: N/A AZD1222 (v1) 7d			p4	F	66	PMH: N/A	AZD1222 (v1)	7d
FH: N/A				_	•	FH: N/A		
p9 F 39 first clinical CIS episode in 2019 with a complete recovery, FH: N/A BNT162b2 (v1) 3d	— 11 6 1	a	p9	F	39	first clinical CIS episode in 2019 with a complete recovery, FH: N/A	BNT162b2 (v1)	3d
Tagliaferri Case report I F 32 PMH: Neg BNT162b2 (v1) /d	Tagliaterri	Case report	1	F	32	PMH: Neg	BNT162b2 (v1)	7d
FH: N/A	et al. ⁴⁰	<u> </u>	4		20	FH: N/A		4.1
Toljan et al. ²² Case series p1 F 29 PMH: migraines BNT162b2 (v1) 1d	Toljan et al. ²²	Case series	pI	F	29	PMH: migraines	BNT162b2 (v1)	Id
(n = 5)		(n = 5)	2	м	27	FH: N/A		1
p2 M 37 PMH: Neg Initiated: BNT162b2 v1			p2	М	37	PMH: Neg	Initiated: BNT162b2	
$FH: N/A \qquad (VI), Develope: 3d$						FH: N/A	(VI), Develope:	30
BN116262 (V2) V2							BN116262 (v2)	v2
5W mDNA 1272 (v2)			-2	м	41	DMIL Nog	$mDN(\Lambda 1072 (m2))$	3W
$\begin{array}{cccccccccccccccccccccccccccccccccccc$			ps	IVI	41	FILL N/A	$\operatorname{IIIKINA-1275}(V2)$	~ 1
ΓΠ: Ν/Α III n4 E 46 unileteral entic neuritic et 29 preujously unremarkable brein MDI Initiated: mDNA v1;			n /	Б	16	ΓΠ: N/A unilataral ontio nauritic at 29, proviously unromarkable brain MPI	Initiated: mDNA	1111
p^{4} r^{5} 40 unmaterial optic neutrits at 36, previously unrematikable brain wiki initiated. Initiate			P4	I.	40		1273 (y1)	VI.
111. 10/A 12/3 (v1), 10/A Developed: v2:						111.10/R	1273(V1), Developed:	$\frac{1N}{A}$
$mRN\Delta_{-}1273 (y2) = 3d$							mRNA_1273 (v2)	v2. 3d
n5 E 43 N/A BNT162b2 (v2) 5w			n5	F	43	N/Δ	BNT162b2 (v2)	5w
Watad et al ³⁰ Case series ^{***} p7 F 45 PMH: Controlled hypothyroidism BNT16262 (v2) $7d$	Watad et al ³⁰	Case series***	p5 n7	F	45	PMH: Controlled hypothyroidism	BNT162b2 (v2) BNT162b2 (v1)	7d
(n = 27)	watad of al.	(n = 27)	P'		15	FH: N/A	BI(110202 ((1)	74
Kim et al 42 Case series $^{\ddagger }$ p1 E 28 PMH: Neg BNT162b2 (v2) 28d	Kim et al ⁴²	Case series ^{‡‡}	n1	F	28	PMH: Neg	BNT162b2 (v2)	28d
(n = 10) FH: N/A	Thin of uit	(n = 10)	P	-	20	FH: N/A	BI(110202 (12)	200
p2 F 29 PMH: thyroid cancer BNT162b2 (v2) 8d		(p2	F	29	PMH: thyroid cancer	BNT162b2 (v2)	8d
FH: N/A			r-	-	_,	FH: N/A	(`_)	
Ebrahimi et al. ⁴⁶ Case series p1 M 24 PMH: Headache BBIBP-CorV (v2) 14d	Ebrahimi et al.46	Case series	p1	М	24	PMH: Headache	BBIBP-CorV (v2)	14d
(n = 12) FH: Neg		(n = 12)	T,			FH: Neg	· · · · · · · · · · · · · · · · · · ·	
p2 F 46 PMH: OCD, IDA, depression BBIBP-CorV (v2) 3d		× /	p2	F	46	PMH: OCD, IDA, depression	BBIBP-CorV (v2)	3d
FH: Neg			r			FH: Neg	· · /	

Table 3. Part I: Studies specifically addressing the new diagnosis of MS following SARS-CoV-2 vaccination

MS onset following Sinopharm COVID-19 vaccination

Study	Design	Ν	Sex	Age (year)	PMH, FH	Vaccine type (dose)	Int.
Ebrahimi et al.46	Case series	p3	F	42	PMH: hyperthyroidism, hypertension, Asthma	BBIBP-CorV (v1)	20d
	(n = 12)				FH: Neg		
		p4	F	21	PMH: IDA	BBIBP-CorV (v2)	2d
					FH: MS in mother and cousin		
		p5	F	20	PMH: Neg	BBIBP-CorV (V2)	60d
					FH: Neg		
		рб	F	23	PMH: Neg	BBIBP-CorV (v1)	10d
					FH: MS in sister		
		p7	F	19	PMH: IDA, hypothyroidism, OCD	BBIBP-CorV (v2)	10d
					FH: Neg		
		p8	F	50	PMH: OCD	AZD1222 (v2)	11d
					FH: Neg		
		p9	F	30	PMH: Neg	BBIBP-CorV (v3)	6d
					FH: Neg		

Table 3. Part I: Studies specifically addressing the new diagnosis of MS following SARS-CoV-2 vaccination (continue)

Table 3. Part II: Studies specifically addressing the new diagnosis of MS following SARS-CoV-2 vaccination

Study	Initial symptoms, signs	EDSS	MRI	Labs and CSF	Treat., response	Notes
				analyses	_	
Fujimori et al. ²⁵	Rt. hand: numbness and sensory imp, ascended to Rt. Shoulder, Rt. cervical 5 th -8 th dermatome sensory imp	N/A	Brain: several PV or SC T2 hyperintense white matter lesions w/o Gad enhancement no brainstem lesions Cervical: a T2 hyperintense right spinal cord lesion with Gad enhancement at C5/C6 Thoracic: N/A	mildly elevated leukocytes protein and glucose: NL, OCB: pos, IgG index: 1.04, MBP < 102 pg/ml, IL-6 < 4.0 pg/ml, MOG-IgG: Neg	IVMP (3 g) recovered	The patient possibly had pre-existing subclinical inflammatory CNS disease before vaccination, since the patient already had several asymptomatic non-gadolinium enhanced brain lesions and oligoclonal IgG band on admission. Although she had a history of left peripheral facial nerve palsy that resolved after steroid therapy, we could not confirm the episode as her initial clinical manifestation of MS.
Havla et al. ²⁴	Lt. abdominal neuropathic pain sensory imp below the T6 level, Rt. abdominal wall and genital hypoesthesia, Lt. leg paresis	N/A	Brain: > 20 partially confluent lesions with spatial dissemination w/o Gad enhancement Cervical: NL Thoracic: a contrast-enhancing lesion at T6	Mild pleocytosis, OCB: pos, IgG index: N/A	IVMP (1 st cycle: 5 g, 2 nd cycle:10 g), TPE (No complete remission [*])	Assuming that some of these vaccines do carry a small risk of autoimmune exacerbation, it is still unclear whether and how this might differ between the different vaccines and whether patients with pre- existing inflammatory CNS disease should be prioritized for any particular vaccine.
Khayat-Khoei et al. ²³	Rt. eye visual symptoms mild blurring, progressed over the next few days pain with eye movement OD, relative RAPD, decreased visual acuity, color desaturation OD, monocular central/inferior monocular deficit	0#	Brain: multiple (9) T2 hyperintense, PV, SC, posterior fossa lesions, with one Gad enhancement Spinal cord: multiple (> 2) T2 lesion, with one Gad-enhanced lesion	IgG index: 1.27, elevated cell count, protein and glucose: NL, OCB: Neg	IVMP (5 g) recovered	Our report is anecdotal and does not prove a cause-and-effect relationship between SARS-CoV-2 mRNA vaccines and active CNS demyelinating disease. We do not know the number of people with MS who were vaccinated against COVID-19 in the communities from which these cases were derived.
	Unilateral painless vision blurring, visual acuity (20/50 OS)	0	Brain: multiple (7) T2 hyperintense white matter lesions with a single gadolinium-enhancing lesion Spinal cord: one new T2 lesion, w/o Gad-enhancement	OCB: pos (> 5), elevated IgG index, normal CSF cell count, protein, and glucose. NMO antibody: Neg	IVMP (3 g) recovered	
Mathew and John ⁴¹	Lt. upper and lower limbs paresthesia, Lhermitte's sign	N/A	lesions in the brain and spinal cord (not described), Two lesions in the brain and one in the spinal cord were enhancing	N/A	IVMP (5 g) partially improved ^{**}	She was diagnosed with MS in view of two episodes of neurological dysfunction four years apart (Dissemination in time) and multiple lesions on MRI in the brain and spinal cord (Dissemination in space) with a probable vaccine induced second relapse.

MS onset following Sinopharm COVID-19 vaccination

Table 3. Part II: Studies specifically addressing the new diagnosis of MS following SARS-CoV-2 vaccination (continue)

Study	Initial symptoms, signs	EDSS	MRI	Labs and CSF	Treat., response	Notes
				analyses		
Nistri et al. ⁴³	Rt. eye visual acuity deficit	2.0	Brain: enhancing lesion in the corpus callosum, multiple white matter unenhanced lesions, and lesions in the occipital lobe Spine: NL	N/A	IVMP (N/A) marked improvement	Although the evidence of an association between vaccination and MS activity is still debated, a link between them has been suggested, within the first 30 days after immunization, given the possibility that
	Visual disturbance, Rt. sided postural instability	2.5	Brain: multiple white matter lesions, four of them enhancing in the left para-trigonal and PV white matter Spine: NL	OCB: pos	IVMP (N/A) partial improvement	vaccines may accelerate the transition from subclinical to clinical disease through a stimulation of the immune system.
	Rt. limbs dysesthesia	1.0	Brain: a new enhancing lesion in the mesencephalon Spine: NL	N/A	IVMP (N/A) good recovery	
Tagliaferri et al. ⁴⁰	Rt. sided weakness, Rt. hand fine motor weakness, word slurring, gait instability, diffuse Rt. sided weakness, Rt. upper and lower EXT diminished strength and sensation	N/A	Brain: multiple round hyperintensities in the white matter with restricted diffusion in the left pons Cervical: N/A Thoracic: N/A	elevated myelin basic protein, OCB: pos (> 6), IgG index: N/A	IVMP (3 g) response: N/A	Although it remains completely unclear, we associated the MS incidence with the vaccine based on the temporal relationship between receiving the vaccine and onset of symptoms. The purpose of this paper is not to definitely associate the COVID vaccine with the disease; rather, we aim to shed light on the possibility of this rare occurrence.
Toljan et al. ²²	Lt. leg acute onset weakness and numbness, Rt. arm paresthesia, Lt. arm weakness, orbiting sign, mild pronator drift, Lt. leg weakness, Lt. EXT marked hyperreflexia, Lt. leg: diminished vibratory sensation	N/A	Brain: multiple brain lesions, including several PV and JC white matter lesions with one enhancing lesion in the right centrum semiovale Cervical: NL Thoracic: N/A	Pleocytosis, IgG index: 0.71, OCB: pos (10)	IVMP (5 g) significant improvement	The association cannot be determined to be causal, as latent CNS demyelinating disease may unmask itself in the setting of an infection or a systemic inflammatory response.
	Lt. hand: paresthesia, paresthesia spread over the entire Lt. arm, urinary urgency, gait imbalance, Rt. sided internuclear ophthalmoplegia, Lt. arm isolated hyperreflexia	N/A	Brain: multiple PV non- enhancing T2/FLAIR hyperintensities Cervical: a C3-C4 cord T2 and STIR hyperintense lesion Thoracic: N/A	OCB: N/A, IgG index: N/A, Serum AQP4- IgG: Neg, MOG-IgG: Neg	High dose oral PSL, taper over 12 days, 3 days of 600 mg oral PSL, response: N/A	
	Bilateral foot numbness, progressive paraparesis, difficulty initiating voiding, Rt. Hemiparesis, Rt. facial droop	N/A	Brain: multiple intracranial PV and JC T2/FLAIR hyperintensities, with most lesions demonstrating contrast enhancement	Pleocytosis, IgG index: 5.82, OCB: pos (6), Serum AQP4-IgG: NegMOG-IgG: Neg	IVMP (5 g), TPE (5 sess) Oral PSL, No complete remission [¥]	

Table 3. Part II: Studies specifically addressing the new diagnosis of MS following SARS-CoV-2 vaccination (continue)

Study	Initial symptoms, signs	EDSS	MRI	Labs and CSF analyses	Treat., response	Notes
Toljan et al. ²²	Rt. leg intermittent numbness, bilateral arm pain, Lt. lateral abdomen burning sensation, Lt. foot drop, mild anisocoria, Rt. eye red desaturation w/o RAPD, symmetrically reduced strength in lower EXT, diffuse	N/A	Complete spinal cord: additional multifocal enhancing and non- enhancing T2 and STIR dorsal cord hyperintensities Brain: PV and JC intracranial lesions with enhancement of the PV lesion Cervical and thoracic spine: multiple enhancing lesions in the spine	index: 1.83, OCB: pos (20), herpes viruses (HSV, VZV, EBV, CMV), WNV, Borrelia burgdorferi IgG and IgM: Neg VDRL: Neg	IVMP (5 g) response: N/A	
	hyperreflexia with bilateral extensor response distal Rt. arm weakness, Rt.	N/A	Brain: enhancing and non-	MOG-IgG: Neg AQP4-IgG: Neg OCB: pos (8), CSF cell	IVMP (3 g),	
	periorbital and palatal numbness		enhancing temporal and callosal PV ovoid lesions, enhancement of the proximal Rt. trigeminal nerve Cervical: NL Thoracic: NL	counts: NL	Symptoms, Improved	
Watad et al. ³⁰	Lt. leg weakness, Disequilibrium, lower limbs distal numbness	N/A	Brain: Multiple PV white matter changes Cervical: N/A Thoracic: N/A	OCB: pos IgG index: N/A	IVPM (5 g), Rapid improvement	Most of the reported disease were flares, which supports the idea of the delicate balance of immune homeostasis in such cases being momentarily tipped into a pro- inflammatory state by vaccination.
Kim et al. ⁴²	Unilateral optic neuritis	N/A	Orbit: Rt. optic nerve T2 HSI with enhancement Brain: Multiple T2 HSI on PV, brainstem with enhancement Spine: NL	WBC: 5, OCB: pos, IgG index: 0.48	IVMP (3 g), Complete recovery	There is no definitive way to link the onset of CNS-IDDs with COVD- 19 vaccination, but the close temporal association may suggest a pathogenic link.
	Unilateral optic neuritis	N/A	Orbit: Lt. optic nerve T2 HSI with enhancement Brain: Multiple T2 HSI on PV, JC with enhancement Spine: ss myelitis (T11-12 level)			
Ebrahimi	Balance disturbance	3	N/A	N/A	IVMP (3 g),	We see both old and new lesions
et al. ⁴⁶		2.5	N 7/A	NT / A	marked improvement	meaning that this excessive immune
	Rt. hand paresthesia	2.5	N/A	N/A	IVMP (3 g), nartial response ^{###}	of these diseases following vaccination in
	Lt. hand paresthesia to complete numbness	2	N/A	N/A	IVMP (3 g), complete recovery	those who already have a weakened immune system.

Table 3. Part II: Studies spe	cifically addressing the new	diagnosis of MS following	g SARS-CoV-2 vaccination (continue)

Study	Initial symptoms, signs	EDSS	MRI	Labs and CSF	Treat., response	Notes
				analyses		
Ebrahimi	Lt. hand paresthesia	N/A	N/A	N/A	IVMP (3 g), partial	
et al. ⁴⁶					response ^{¥¥}	
	Lt. hand paresthesia, Lhermitte	2.5	N/A	N/A	IVMP (3 g), partial	
	sign				response	
	Rt. sided facial paresthesia, Rt.	1.5	N/A	N/A	IVMP (3 g),	
	hand tingling				complete recovery	
	Lt. sided numbness	1.5	N/A	N/A	IVMP (3 g),	
					complete recovery	
	Numbness in Lt. hand and foot	2.5	N/A	N/A	IVMP (3 g), EDSS	
					improved	

AQP4-IgG: Aquaporin 4-IgG; CIS: Clinically isolated syndrome; CMV: Cytomegalovirus; EBV: Epstein–Barr virus; EXT: Extremities; FH: Family history; g: Gram; HIS: High signal intensity; HSV: Herpes-simplex virus; IDA: Iron deficiency anemia; IMP: Impairment; INT: Interval; m: Month; JC: Juxtacortical; MBP: Myelin basic protein; MOG-IgG: Myelin oligodendrocyte glycoprotein-IgG; MS: Multiple sclerosis; Neg: Negative; NL: Normal; Numb.: Numbness; OCB: Oligoclonal band; OCD: Obsessive-compulsive disorder; pos: Positive; PMH: Past medical history; PSL: Prednisolone; PV: Periventricular; RAPD: Relative afferent pupillary defect; SC: Subcortical; sess: session(s); ss: Short-segment; STIR: Short-tau inversion recovery; Treat.: Treatment; v1: 1st vaccine dose; v2: 2nd vaccine dose; VZV: Varicella-zoster virus; WNV: West Nile virus; w/o: Without; CNS: Central nervous system

*Complete remission of symptoms did not occur even after a second cycle of glucocorticoid therapy (2000 mg IVMP for 5 days) and escalating the relapse therapy with plasma exchange treatment was performed, which resulted in further improvement.

**At six weeks she had improved 80% with mild distal paresthesia and persisting Lhermitte symptom.

***In this case series, 27 patients with new-onset or flare of immune-mediated diseases were included, one of which was diagnosed with new-onset multiple sclerosis (patient 7).

[‡]In this series, 7 patients who developed neurologic symptoms and MRI findings consistent with active CNS demyelination of the optic nerve, brain, and/or spinal cord were reported. The final diagnosis was exacerbation of known stable MS (n = 4, two were receiving disease-modifying therapy (DMT) at the time of vaccination), new onset MS (n = 2), or new onset neuromyelitis optica (NMO) (n = 1). In this table only data for the 2 patients with new onset MS is reported (patients 2 and 5).

^{‡‡}In this study, among 117 cases, 10 had their first disease manifestation within one month following COVID-19 vaccination, two of which were diagnosed with MS.

****In this case series, 12 patients developing MS, clinically isolated syndrome (CIS), and NMOSD following COVID 19 vaccines were reported, 9 of which were diagnosed with MS (patients 1 to 9).

[#]At baseline and after treatment

##In this study, 16 patients received a diagnosis of MS, three of which had a first episode after COVID-19 vaccination (patients 2, 4, and 9).

###Forty-two days after her diagnosis, numbness was resolved, but she experienced frequent urination and urinary incontinence one to three times a week. After a four-month follow-up, she said that her urinary dysfunctions were no longer an issue and that she was seldom bothered by them.

⁴Residual Rt. upper extremity weakness and symmetric hyperreflexia with sustained clonus in both ankles, residual decreased sensation to all sensory modalities over the entire Lt. arm, left half of trunk below T4 level dermatome, and over the entire Lt. leg.

"Mild right wrist extensor weakness and ipsilateral hip flexor and knee flexor weakness persist after a week.

^{¥¥}She feels no numbness in her fingers anymore, but as she bends her neck, she still feels some trembling in her back.

After two months, she still feels a light electric shock passing down her neck, but her numbness and tingling have gone away.

Remarkably, 6/26 patients had a previous history of neurological conditions, including headaches/migraine (n = 2),^{22,46} recovered facial palsy (n = 1),²⁵ recovered facial numbness with left upper limb weakness (n = 1),⁴¹ unilateral optic neuritis with previously unremarkable brain MRI (n = 1),²² and the first episode of the clinically isolated syndrome (CIS) with complete recovery (n = 1).⁴³ Only 2/7 studies, accounting for 10 individuals, provided information about the patient's family history.^{24,46} While family history was unremarkable in 7/10, 3/10 patients had positive family histories, with MS present in a paternal cousin,²⁴ in both mother and cousin,⁴⁶ and in a sister.⁴⁶

MS symptoms manifested within 1 day to 3 months after receiving BNT162b2 (n = 11; v1: 6, v2: 5), $^{22-25,30,40,42,43}$ mRNA-1273 (n = 3; v1: 1, v2: 2),^{22,23} BBIBP-CorV (n = 8; v1: 2, v2: 5, v3: 1),⁴⁶ or AZD1222 (n = 4, v1: 2, v2: 2).41,43,46 The duration between the symptoms onset and vaccination was less than 6 weeks in 24/26 patients (≤ 7 days in 12/26), more than 6 weeks in 1/21 (p5 in the study by Ebrahimi et al.),46 and remained unspecified in 1/21 (p4 in the study by Toljan et al.).²² It is noteworthy that 2/21 patients (p2 and p4 in the study by Toljan et al.) experienced MS-related symptoms after both vaccine doses.²² One patient (p2) experienced initial symptoms 3 days after receiving v1, with further progression observed 3 weeks after the subsequent dose.²² While the authors did not specify the duration between the other patient's (p4) initial symptoms and v1 administration, they did note that the symptoms, which had previously commenced, progressed 3 days after the administration of v2.22

Brain MRI results were available in all studies except one.46 In line with our findings, periventricular lesions were the most frequently lesions, 22, 23, 25, 30, 42, 43 observed followed bv subcortical^{23,25} and juxtacortical lesions.^{22,42} Spinal cord MRI findings were reported in all studies except 3,30,40,46 with lesions detected at C3-C4,22 C5-C6,²⁵ T6,²⁴ and T11-T12.⁴² MS-related laboratory evaluation results were reported for all patients except 3.41,43,46 In most cases, the diagnosis of MS was supported by CSF pleocytosis, positive OCBs, and elevated IgG index. All patients received 3-5 gr of IVMP, except for 1 patient (p2 in the study by Toljan et al.), who received high-dose oral prednisolone.²² Additionally, 2/26 patients underwent TPE.22,24 At the last follow-up, all patients showed partial to complete symptom recovery. Response to treatment was not reported for 3 patients.^{22,40}

Although the exact pathophysiology behind some autoimmune sequelae in the context of vaccine exposure is currently unknown, there is a hypothesis that COVID-19 vaccines might trigger an excessive inflammatory immune reaction in a subgroup of vulnerable individuals. This could potentially accelerate the transition from subclinical to clinical disease and reveal a previously concealed demyelinating condition within the CNS,³² which is not surprising due to the involvement of neuroinflammation in various neurological conditions such as MS.64,65 Notably, a combination of old and new MRI lesions in many of the reported patients could indicate a clinically latent disease, which was masked before the vaccination-induced immune response.22-25 Although only 1 of our patients showed evidence of a latent demyelinating process, we cannot exclude possible genetic Other susceptibility in others. suggested mechanisms include molecular mimicry (similarities between self-antigens and vaccine), aberrant immune response, vaccine-related factors (i.e., adjuvants), and an already altered immune response in susceptible individuals.28

Determining whether these observations imply causality (the development of a new disease) or mere temporal coincidence (the manifestation of an existing, previously subclinical neuroinflammation) requires thorough consideration of various issues. Firstly, although there have been reports of autoimmune diseases occurring after vaccination, only a limited number have been definitively labeled as "vaccine-induced".66,67 Butler et al. have proposed specific criteria for establishing "probable causality" in cases of neurological AEFI, including a typical time frame (< 6 weeks), no indication of an alternative etiology, and no other risk factors⁵⁰ (Table 1). Secondly, even when considering the possibility of a small risk of autoimmune exacerbation associated with SARS-CoV-2 vaccines, research has indicated a substantially higher risk of neurological AEs, including CNS demyelinating disease, following infections as opposed to vaccination.27,28,68 Eventually, our understanding of this will only be enhanced when forthcoming studies investigate whether COVID-19 vaccines raise the likelihood of MS beyond the anticipated, usual background rate of such incidents.

Conclusion

This study is the second case series to report MS

incidence following the administration of BBIBP-CorV in a temporal relationship. Apart from our patients, we conducted a thorough review of the relevant published literature. Among our 8 patients, 1 case was deemed unlikely to be vaccine-related due to a prior history of facial numbness together with brain atrophy and black holes in MRI. Nevertheless, an association of < 6 weeks with ictus and evidence of OCBs along with the MRI findings suggests the possibility of a latent disease that may have been triggered by vaccination. It is worth mentioning that the occurrence of MS onset following vaccination is still extremely rare when compared to the vaccinated population. As with previous reports, due to their descriptive nature, lack of a comparison group, and limited sample, it is not possible to establish a causal relationship between vaccination and any specific AEFI based on single case reports;69 hence, the findings should be interpreted with caution. Causality can change when additional information about the same or similar cases becomes available. Although the short interval between the exposure and event might reduce the possible role of potentially confounding factors, several steps should be taken to establish a causal relationship beyond a mere temporal association between exposure and the event. Strength of the association, dose-response relationship (higher likelihood of the outcome with exposures), replication, biological higher plausibility, coherence (consistency of the association with existing knowledge), and analogy are among the most important criteria to establish a causal relationship.70 Statistical analyses also play a crucial role in establishing causality in research by identifying correlations, estimating the strength of the association, and controlling for confounding variables.⁷⁰ Collectively, the current evidence strongly suggests that the benefits of vaccination outweigh the plausible risks.

Conflict of Interests

The authors declare no conflict of interest in this study.

Acknowledgments

We thank all patients and the staff in our center for their kind assistance.

This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors.

References

- Hu B, Guo H, Zhou P, Shi ZL. Characteristics of SARS-CoV-2 and COVID-19. Nat Rev Microbiol 2021; 19(3): 141-54.
- Cucinotta D, Vanelli M. WHO declares COVID-19 a pandemic. Acta Biomed 2020; 91(1): 157-60.
- World Health Organization. WHO Coronavirus (COVID-19) Dashboard [Online]. [cited 2022 Feb 28]; Available from: URL: https://covid19.who.int/
- Zali A, Khodadoost M, Gholamzadeh S, Janbazi S, Piri H, Taraghikhah N, et al. Mortality among hospitalized COVID-19 patients during surges of SARS-CoV-2 alpha (B.1.1.7) and delta (B.1.617.2) variants. Sci Rep 2022; 12(1): 18918.
- Jameie M, Safarian NM, Mansouri P, Jalali A, Aghajani F, Lotfi-Tokaldany M, et al. The impact of the COVID-19 pandemic on hospitalization rates due to prosthetic valve thrombosis. J Tehran Heart Cent 2023; 18(2): 136-41.
- Soheili A, Khani S, Montazeri S, Shayegh A, Miragha M, Jameie M, et al. COVID-19 and acute kidney injury presentation; stages and prognosis. J Prev Epidemiol 2021; 6(1): e15.
- Majidi F, Mohagheghi Dare RA, Jameie M, Jameie M, Mansouri P, Varpaei HA, et al. The relationship between cardiological parameters and PCR in patients with coronavirus infection: A cross-sectional

study. Medicine (Baltimore) 2022; 101(50): e31935.

- Wang C, Tee M, Roy AE, Fardin MA, Srichokchatchawan W, Habib HA, et al. The impact of COVID-19 pandemic on physical and mental health of Asians: A study of seven middle-income countries in Asia. PLoS One 2021; 16(2): e0246824.
- Duden GS, Reiter J, Paswerg A, Weibelzahl S. Mental health of healthcare professionals during the ongoing COVID-19 pandemic: A comparative investigation from the first and second pandemic years. BMJ Open 2023; 13(3): e067244.
- Borhany H, Golbabaei S, Jameie M, Borhani K. Moral decision-making in healthcare and medical professions during the COVID-19 pandemic. Trends in Psychol 2023; 31(1): 210-30.
- Chen X, Laurent S, Onur OA, Kleineberg NN, Fink GR, Schweitzer F, et al. A systematic review of neurological symptoms and complications of COVID-19. J Neurol 2021; 268(2): 392-402.
- World Health Organization. Coronavirus Disease (COVID-19): Vaccines [Online]. [cited 2022 Mar 1]; Available from: URL: https://www.who.int/news-room/questionsand-answers/item/coronavirus-disease-(covid-19)-

vaccines?gclid=CjwKCAiAgvKQBhBbEi wAaPQw3GLluFkuSSoT_AlhcXDldrPzX pnUkkciDrC70HeFHja3PHifQkQ5MxoC AoIQAvD_BwE&topicsurvey=v8kj13

- Walton C, King R, Rechtman L, Kaye W, Leray E, Marrie RA, et al. Rising prevalence of multiple sclerosis worldwide: Insights from the Atlas of MS, third edition. Mult Scler 2020; 26(14): 1816-21.
- Vandebergh M, Degryse N, Dubois B, Goris A. Environmental risk factors in multiple sclerosis: Bridging Mendelian randomization and observational studies. J Neurol 2022; 269(8): 4565-74.
- Abrahamyan S, Eberspacher B, Hoshi MM, Aly L, Luessi F, Groppa S, et al. Complete Epstein-Barr virus seropositivity in a large cohort of patients with early multiple sclerosis. J Neurol Neurosurg Psychiatry 2020; 91(7): 681-6.
- Najafi S, Ghane M, Poortahmasebi V, Jazayeri SM, Yousefzadeh-Chabok S. Prevalence of cytomegalovirus in patients with multiple sclerosis: A case-control study in northern Iran. Jundishapur J Microbiol 2016; 9(7): e36582.
- 17. Rice EM, Thakolwiboon S, Avila M. Geographic heterogeneity in the association of varicella-zoster virus seropositivity and multiple sclerosis: A systematic review and meta-analysis. Mult Scler Relat Disord 2021; 53: 103024.
- Kazmierski R, Wender M, Guzik P, Zielonka D. Association of influenza incidence with multiple sclerosis onset. Folia Neuropathol 2004; 42(1): 19-23.

- Khan F, Sharma P, Pandey S, Sharma D, Vijayavarman V, Kumar N, et al. COVID-19-associated Guillain-Barre syndrome: Postinfectious alone or neuroinvasive too? J Med Virol 2021; 93(10): 6045-9.
- Mahapure KS, Prabhune AS, Chouvhan AV. COVID-19-associated acute disseminated encephalomyelitis: A systematic review. Asian J Neurosurg 2021; 16(3): 457-69.
- Valiuddin H, Skwirsk B, Paz-Arabo P. Acute transverse myelitis associated with SARS-CoV-2: A case-report. Brain Behav Immun Health 2020; 5: 100091.
- Toljan K, Amin M, Kunchok A, Ontaneda D. New diagnosis of multiple sclerosis in the setting of mRNA COVID-19 vaccine exposure. J Neuroimmunol 2022; 362: 577785.
- Khayat-Khoei M, Bhattacharyya S, Katz J, Harrison D, Tauhid S, Bruso P, et al. COVID-19 mRNA vaccination leading to CNS inflammation: A case series. J Neurol 2022; 269(3): 1093-106.
- Havla J, Schultz Y, Zimmermann H, Hohlfeld R, Danek A, Kumpfel T. First manifestation of multiple sclerosis after immunization with the Pfizer-BioNTech COVID-19 vaccine. J Neurol 2022; 269(1): 55-8.
- Fujimori J, Miyazawa K, Nakashima I. Initial clinical manifestation of multiple sclerosis after immunization with the Pfizer-BioNTech COVID-19 vaccine. J Neuroimmunol 2021; 361: 577755.
- Goss AL, Samudralwar RD, Das RR, Nath A. ANA investigates: Neurological complications of COVID-19 vaccines. Ann Neurol 2021; 89(5): 856-7.
- Patone M, Handunnetthi L, Saatci D, Pan J, Katikireddi SV, Razvi S, et al. Neurological complications after first dose of COVID-19 vaccines and SARS-CoV-2 infection. Nat Med 2021; 27(12): 2144-53.
- Ismail II, Salama S. A systematic review of cases of CNS demyelination following COVID-19 vaccination. J Neuroimmunol 2022; 362: 577765.
- Alroughani R, Al-Hashel J, Abokalawa F, AlMojel M, Farouk AS. COVID-19 vaccination in people with multiple sclerosis, real-life experience. Clin Neurol Neurosurg 2022; 220: 107374.
- 30. Watad A, De Marco G, Mahajna H, Druyan A, Eltity M, Hijazi N, et al. Immune-mediated disease flares or new-onset disease in 27 subjects following mRNA/DNA SARS-CoV-2 vaccination. Vaccines (Basel) 2021; 9(5): 435.
- Farez MF, Correale J. Immunizations and risk of multiple sclerosis: Systematic review and meta-analysis. J Neurol 2011; 258(7): 1197-206.
- 32. Langer-Gould A, Qian L, Tartof SY, Brara SM, Jacobsen SJ, Beaber BE, et al. Vaccines and the risk of multiple sclerosis and other central nervous system demyelinating diseases. JAMA Neurol 2014; 71(12): 1506-13.
- Scheller NM, Svanstrom H, Pasternak B, Arnheim-Dahlstrom L, Sundstrom K, Fink K, et al. Quadrivalent HPV vaccination

and risk of multiple sclerosis and other demyelinating diseases of the central nervous system. JAMA 2015; 313(1): 54-61.

- Mailand MT, Frederiksen JL. Vaccines and multiple sclerosis: A systematic review. J Neurol 2017; 264(6): 1035-50.
- Hapfelmeier A, Gasperi C, Donnachie E, Hemmer B. A large case-control study on vaccination as risk factor for multiple sclerosis. Neurology 2019; 93(9): e908-e916.
- 36. Huttner A, Eperon G, Lascano AM, Roth S, Schwob JM, Siegrist CA, et al. Risk of MS relapse after yellow fever vaccination: A self-controlled case series. Neurol Neuroimmunol Neuroinflamm 2020; 7(4): e726.
- 37. Jameie M, Togha M, Looha MA, Hemmati N, Jafari E, Nasergivehchi S, et al. Clinical characteristics and factors associated with headaches following COVID-19 vaccination: A cross-sectional cohort study (P14-12.003). Neurology 2023; 100 (17_Suppl_2): 4105.
- Garcia-Grimshaw M, Ceballos-Liceaga SE, Hernandez-Vanegas LE, Nunez I, Hernandez-Valdivia N, Carrillo-Garcia DA, et al. Neurologic adverse events among 704,003 first-dose recipients of the BNT162b2 mRNA COVID-19 vaccine in Mexico: A nationwide descriptive study. Clin Immunol 2021; 229: 108786.
- 39. VAERS Team. Vaccine Adverse Event Reporting System (VAERS) Standard Operating Procedures for COVID-19 (as of 29 January 2021) [Online]. [cited 2023 Sep 4]; Available from: URL: https://www.cdc.gov/vaccinesafety/pdf/V AERS-v2-SOP.pdf
- Tagliaferri AR, Horani G, Stephens K, Michael P. A rare presentation of undiagnosed multiple sclerosis after the COVID-19 vaccine. J Community Hosp Intern Med Perspect 2021; 11(6): 772-5.
- Mathew T, John SK. COVID-19 vaccine (ChAdOx1 nCoV-19 Corona virus vaccine (Recombinant) – COVISHIELD related MS relapse. Neuroimmunology Reports 2021; 1: 100006.
- 42. Kim KH, Kim SH, Park NY, Hyun JW, Kim HJ. Onset of various CNS inflammatory demyelination diseases following COVID-19 vaccinations. Mult Scler Relat Disord 2022; 68: 104141.
- 43. Nistri R, Barbuti E, Rinaldi V, Tufano L, Pozzilli V, Ianniello A, et al. Case report: Multiple sclerosis relapses after vaccination against SARS-CoV2: A series of clinical cases. Front Neurol 2021; 12: 765954.
- 44. Nabizadeh F, Ramezannezhad E, Kazemzadeh K, Khalili E, Ghaffary EM, Mirmosayyeb O. Multiple sclerosis relapse after COVID-19 vaccination: A case report-based systematic review. J Clin Neurosci 2022; 104: 118-25.
- 45. Etemadifar M, Abhari AP, Nouri H, Sigari AA, Piran Daliyeh SM, Maracy MR, et al. Self-reported safety of the BBIBP-CorV (Sinopharm) COVID-19 vaccine among Iranian people with multiple sclerosis. Hum Vaccin

Immunother 2022; 18(1): 2041945.

- Ebrahimi N, Mazdak M, Shaygannejad V, Mirmosayyeb O. CNS demyelinating disease following inactivated or viral vector SARS-CoV-2 vaccines: A case series. Vaccine 2023; 41(5): 1003-8.
- 47. Gagnier JJ, Kienle G, Altman DG, Moher D, Sox H, Riley D. The CARE guidelines: Consensus-based clinical case reporting guideline development. J Med Case Rep 2013; 7: 223.
- World Medical Association. World Medical Association Declaration of Helsinki: Ethical principles for medical research involving human subjects. JAMA 2013; 310(20): 2191-4.
- Thompson AJ, Banwell BL, Barkhof F, Carroll WM, Coetzee T, Comi G, et al. Diagnosis of multiple sclerosis: 2017 revisions of the McDonald criteria. Lancet Neurol 2018; 17(2): 162-73.
- Butler M, Tamborska A, Wood GK, Ellul M, Thomas RH, Galea I, et al. Considerations for causality assessment of neurological and neuropsychiatric complications of SARS-CoV-2 vaccines: From cerebral venous sinus thrombosis to functional neurological disorder. J Neurol Neurosurg Psychiatry 2021; 92(11): 1144-51.
- Ramo-Tello C, Blanco Y, Brieva L, Casanova B, Martinez-Caceres E, Ontaneda D, et al. Recommendations for the diagnosis and treatment of multiple sclerosis relapses. J Pers Med 2021; 12(1): 6.
- 52. Mallapaty S. Iran hopes to defeat COVID with home-grown crop of vaccines. Nature 2021; 596(7873): 475.
- 53. World Health Organization. The Sinopharm COVID-19 vaccine: What you need to know [Online]. [cited 2022 Mar 1]; Available from: URL: https://www.who.int/news-room/featurestories/detail/the-sinopharm-covid-19vaccine-what-you-need-to-know
- 54. World Health Organization. Annexes to the interim recommendations for use of the inactivated COVID-19 vaccine BIBP developed by China National Biotec Group (CNBG), Sinopharm, 7 May 2021 [Online]. [cited 2021 May 7]; Available from: URL: https://www.who.int/publications/i/item/W HO-2019-nCoV-vaccines-SAGE_recommendation-BIBP-annexes-2021.1
- 55. Pormohammad A, Zarei M, Ghorbani S, Mohammadi M, Razizadeh MH, Turner DL, et al. Efficacy and safety of COVID-19 vaccines: A systematic review and meta-analysis of randomized clinical trials. Vaccines (Basel) 2021; 9(5): 467.
- Gee J, Marquez P, Su J, Calvert GM, Liu R, Myers T, et al. First month of COVID-19 vaccine safety monitoring - United States, December 14, 2020-January 13, 2021. MMWR Morb Mortal Wkly Rep 2021; 70(8): 283-8.
- 57. Voysey M, Clemens SAC, Madhi SA, Weckx LY, Folegatti PM, Aley PK, et al. Safety and efficacy of the ChAdOx1 nCoV-19 vaccine (AZD1222) against SARS-CoV-2: An interim analysis of four

randomised controlled trials in Brazil, South Africa, and the UK. Lancet 2021; 397(10269): 99-111.

58. Centers for Disease Control and Prevention. People with Certain Medical Conditions [Online]. [cited 2022 Feb 28]; Available from: URL: https://www.cdc.gov/coronavirus/2019ncov/need-extra-precautions/people-withmedical-

conditions.html?CDC_AA_refVal=https %3A%2F%2Fwww.cdc.gov%2Fcoronavi rus%2F2019-

ncov%2Fvaccines%2Frecommendations %2Funderlying-conditions.html

- 59. Jameie M, Togha M, Azizmohammad LM, Jafari E, Yazdan PM, Hemmati N, et al. Characteristics of headaches attributed to SARS-CoV-2 vaccination and factors associated with its frequency and prolongation: A cross-sectional cohort study. Front Neurol 2023; 14: 1214501.
- 60. Abara WE, Gee J, Marquez P, Woo J, Myers TR, DeSantis A, et al. Reports of Guillain-Barre syndrome after COVID-19 vaccination in the United States. JAMA Netw Open 2023; 6(2): e2253845.
- Rafati A, Pasebani Y, Jameie M, Yang Y, Jameie M, Ilkhani S, et al. Association of SARS-CoV-2 vaccination or infection

with bell palsy: A systematic review and meta-analysis. JAMA Otolaryngol Head Neck Surg 2023; 149(6): 493-504.

- 62. Eslait-Olaciregui S, Llinas-Caballero K, Patino-Manjarres D, Urbina-Ariza T, Cediel-Becerra JF, Dominguez-Dominguez CA. Serious neurological adverse events following immunization against SARS-CoV-2: A narrative review of the literature. Ther Adv Drug Saf 2023; 14: 20420986231165674.
- 63. Mirmosayyeb O, Ghaffary EM, Vaheb S, Pourkazemi R, Shaygannejad V. Multiple sclerosis (MS) and neuromyelitis optica spectrum disorder (NMOSD) following COVID-19 vaccines: A systematic review. Rev Neurol (Paris) 2023; 179(4): 265-81.
- 64. Amanollahi M, Jameie M, Heidari A, Rezaei N. The dialogue between neuroinflammation and adult neurogenesis: Mechanisms involved and alterations in neurological diseases. Mol Neurobiol 2023; 60(2): 923-59.
- 65. Amanollahi M, Jameie M, Rezaei N. Neuroinflammation as a potential therapeutic target in neuroimmunological diseases. In: Rezaei N, Yazdanpanah N, editors. Translational Neuroimmunology, Volume 7. 1st ed. Cambridge, MA: Academic Press; 2023. p. 475-504.

- Salemi S, D'Amelio R. Could autoimmunity be induced by vaccination? Int Rev Immunol 2010; 29(3): 247-69.
- 67. Wraith DC, Goldman M, Lambert PH. Vaccination and autoimmune disease: What is the evidence? Lancet 2003; 362(9396): 1659-66.
- 68. Garcia-Grimshaw M, Ceballos-Liceaga SE, Hernandez-Vanegas LE, Nunez I, Hernandez-Valdivia N, Carrillo-Garcia DA, et al. Systemic and neurologic adverse events among 704,003 first-dose recipients of the Pfizer-Biontech (BNT162b2) mRNA COVID-19 vaccine in Mexico [Online]. [cited 2021]; Available from: URL: https://ssrn.com/abstract=3816489 or http://dx.doi.org/10.2139/ssrn.3816489
- World Health Organization. Causality assessment of an adverse event following immunization (AEFI): User manual for the revised WHO classification. Geneva, Switzerland: WHO; 2018.
- Shimonovich M, Pearce A, Thomson H, Keyes K, Katikireddi SV. Assessing causality in epidemiology: Revisiting Bradford Hill to incorporate developments in causal thinking. Eur J Epidemiol 2021; 36(9): 873-87.