REVIEW ARTICLE





Motor imagery for pain and motor function after spinal cord injury: a systematic review

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Received: 20 March 2019 / Revised: 17 November 2019 / Accepted: 21 November 2019 / Published online: 13 December 2019 © The Author(s), under exclusive licence to International Spinal Cord Society 2019

Abstract

Study design Systematic review.

Objectives To evaluate the therapeutic benefits of motor imagery (MI) for the people with spinal cord injury (SCI). **Setting** International.

Methods We searched electronic bibliographic databases, trial registers, and relevant reference lists. The review included experimental and quasi-experimental study designs as well as observational studies. For the critical appraisal of the 18 studies retrieved (three RCT, seven quasi-RCT, eight observational), we used instruments from the Joanna Briggs Institute. The primary outcome measure was pain. Secondary outcome measures included motor function and neurophysiological parameters. Adverse effects were extracted if reported in the included studies. Because of data heterogeneity, only a qualitative synthesis is offered.

Results The included studies involved 282 patients. In most, results were an improvement in motor function and decreased pain; however, some reported no effect or an increase in pain. Although protocols of MI intervention were heterogeneous, sessions of 8–20 min were used for pain treatments, and of 30–60 min were used for motor function improvement. Neurophysiological measurements showed changes in brain region activation and excitability imposed by SCI, which were partially recovered by MI interventions. No serious adverse effects were reported.

Conclusions High heterogeneity in the SCI population, MI interventions, and outcomes measured makes it difficult to judge the therapeutic effects and best MI intervention protocol, especially for people with SCI with neuropathic pain. Further clinical trials evaluating MI intervention as adjunct therapy for pain in SCI patients are warranted.

Introduction

Pain is a common complication after spinal cord injury (SCI), which can be related to effects of the spinal injury, a SCI-imposed lifestyle, or to pre-existing conditions. The most common pain types are nociceptive and neuropathic pain (NeP) [1–3]. Management of chronic pain after SCI is very challenging [4] and recent reviews conclude that there

Emmanuelle Opsommer emmanuelle.opsommer@hesav.ch is still a lack of evidence for the impact of both pharmacological and non-pharmacological treatments [5].

Recent studies have shown that motor cortex stimulation can be used as one of the non-pharmacological approaches to treat pain [6, 7]. It has been proposed that cortical structures involved in movement control might be reorganized [8, 9] as a consequence of the spinal cord lesion causing a mismatch between motor output and sensory feedback [10]. These changes in turn could lead to the pain experience [11]. Correcting this discord between mental body representation, sensory-motor integration, and nociception may help in chronic pain treatment [12].

Motor imagery (MI) is one of the techniques which could be used for this purpose. It is defined as mental representation of movement without any actual body movement or peripheral muscle activation [13–16]. The brain areas (including motor cortex) active during MI and movement itself are largely overlapping [17]. This could explain the fact that mental movement repetition, especially when

Supplementary information The online version of this article (https://doi.org/10.1038/s41393-019-0390-1) contains supplementary material, which is available to authorized users.

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combined with physical practice, improves motor performance in healthy people [18], athletes [19] as well as in individuals with neurological disorders, including SCI [13, 14, 20–22]. However, the effect of MI interventions on pain remains unclear: some studies showed a reduction [10, 23], some an increase [24] and some no effect on pain [25]. There could be many reasons underlying these discrepancies, such as patient's perception of pain, social stressors, patient's expectation from treatment, or MI methodology itself [12, 26]. Indeed, there are different ways to perform MI. It can be carried out from two perspectives: external (third-person) or internal (first-person). The thirdperson perspective is an imagery where a "person views him- or herself from the perspective of an external observer" (i.e., seeing him/herself performing the imagined movement). It is considered to be mainly visual in nature. The first-person perspective requires the person to imagine "being inside his/her body and experiencing those sensations" as if he/she was performing the movement. Therefore, internal (first-person) imagery may include both visual and kinesthetic components [15 p. 945]. Imagery capacity may differ from person to person and should be tested before performing an MI intervention, especially in people with neurological deficits [8]. For example, it was shown that individuals with SCI have difficulties performing MI from the first-person perspective [27]. Several tools exist to assess imagery ability and one of them, the Kinesthetic and Visual Imagery Questionnaire (KVIQ), is especially adapted for individuals with disabilities [28].

To answer questions on the therapeutic benefits of MI interventions on pain in SCI we performed a preliminary search in several databases. The review by Aikat and Dua [29] discussed the therapeutic potential of MI interventions in SCI, but without a critical appraisal of the evidence and not specifically addressing the aspect of pain. The primary purpose of our systematic review was to provide a scrupulous summary of all available primary research on the therapeutic effects of MI interventions on pain in individuals with SCI. A secondary aim was to investigate effects on motor function recovery. Where possible, we extracted information about neurophysiological changes associated with brain activity during MI and discussed the details of MI protocols (i.e., frequency, intensity, duration) for both pain and motor recovery treatments in patients with SCI.

Methods

We published a protocol prior to undertaking this review [12] which was registered with PROSPERO (# CRD42017075144). This systematic review was conducted in accordance with the Joanna Briggs Institute (JBI) methodology for systematic review of effectiveness evidence

[30]. The search strategy, examples of search algorithms with keywords and index terms, as well as information about data extraction procedures, are provided in the Supplementary File 1.

Inclusion and exclusion criteria

This review considered both experimental and quasiexperimental designs as well as observational studies published as full text in English, French, or German (Fig. 1). Studies in other languages were excluded following title appraisal. Studies must have involved primarily adults (18 years and older) with a diagnosis of SCI [12] and have evaluated a MI intervention provided as an independent intervention, added to other therapy, or embedded in therapy. Primary outcomes were those related to pain [31], such as pain intensity (assessed with Visual Analog Scale (VAS), Numerical Rating Scale (NRS), Brief Pain Questionnaire etc.) and pain duration. Secondary outcomes were motor function and activity/disability related outcomes. Additional outcomes were neurophysiological measures of brain activity (i.e., functional magnetic resonance imaging (fMRI), positron emission tomography, electroencephalography (EEG) and magnetoencephalography (MEG)) and motor output (e.g., motor evoked potentials and motor thresholds).

Study selection

A three-step search strategy was used [12]: (1) initial search in PubMed and CINAHL with text word analysis of the title and abstract to identify the keywords and descriptors; (2) secondary search in all databases (JBI Database of Systematic Reviews and Implementation Reports, Cochrane, PubMed, Embase, CINAHL, PsychINFO, PEDro, OTseeker, Campbell, DARE, TRIP, NICE, and BestBets and Bandolier) with identified keywords and index terms; (3) reference lists of all identified reports and articles were searched for additional studies. Individual search strategies were developed for each database to take into account any differences in thesaurus terminology and indexing (example for PubMed in Supplementary file 1). Examples of keywords used: (Spinal Cord Injury OR Spinal Cord Injuries OR Spinal Cord Ischemia OR Paraplegia OR Quadriplegia) AND (Imagery OR Imagination OR Mental Practice OR Cognitive rehearsal OR Guided Imagery OR Motor Imagery). All citations identified were loaded into EndNoteTM and duplicates removed. Titles and then abstracts were screened by two reviewers independently against the inclusion criteria for the review (Fig. 1). The full text of potentially eligible studies was retrieved and assessed against the inclusion criteria by two reviewers independently. Any disagreements that arose between them were resolved through discussion, or with a third reviewer casting a deciding vote.

Fig. 1 PRISMA flow diagram describing screening and review process. *Until 31.01.2019.



Assessment of methodological quality

Two reviewers independently appraised the eligible studies using standardized critical appraisal instruments from the JBI for randomized controlled trials (RCT), quasiexperimental studies, case series, and case reports (https:// joannabriggs.org/critical_appraisal_tools). Any disagreements that arose between the reviewers were resolved through discussion, or with a third reviewer casting a deciding vote. Studies of low methodological quality might bias in the results; therefore, we only included studies of moderate to high quality. Before starting the appraisal, we defined a threshold for each JBI instrument: 6/13 for RCTs, 5/9 for quasi-experimental studies, 6/10 for case series, and 5/8 for case reports.

Data synthesis

Due to heterogeneity across the studies, the findings are presented in narrative form including tables and figures. For the same reason, pain intensity data are displayed in a forest plot with the effect sizes, but without statistical metaanalysis. The effect size was the mean difference between measures of pain intensity. For this, we used Review Manager (RevMan. Version 5.3. Copenhagen: The Nordic Cochrane Centre, The Cochrane Collaboration, 2014).

Results

Figure 1 is the PRISMA flow diagram. Electronic bibliographic databases and additional data source searching returned 10,442 non-duplicate titles. Following title and abstract screening, we screened 117 in full text. Eighteen studies were eligible for the review and critically appraised [10, 11, 20, 21, 23–25, 32–42].

Methodological quality

All studies met the minimum established criteria of moderate to high quality (Supplementary Tables 1–4). Overall, the most frequent risk was selection bias as the majority of the studies did not use randomization for treatment allocation, or did not use a control group. Table 1 describes the population, intervention and comparator as well as the outcomes of the studies included.

Types of studies

Three RCT studies compared two groups of people with SCI [23, 38, 42]. Seven further experimental studies compared either persons with SCI with and/or without NeP to healthy participants, or persons with SCI with NeP to those without NeP [11, 20, 24, 34, 35, 40–42]. Five studies were case series [10, 25, 36, 37, 39] and three were case reports [21, 32, 33].

Participants

The 18 studies involved 282 participants with SCI (mean age: 44.3 ± 11.4 years, range 21–72). The majority of participants with SCI were male (78%). All studies but one [37] (patients in subacute SCI phase with average time after injury of 50 days) included patients with SCI at a chronic stage, with an average time since injury of 7.3 ± 6.1 years (range 0.3-40). The neurological level of injury (NLI) was between C3 and L3 with complete or incomplete lesions. Among individuals with SCI, NeP was present in 166 people and nociceptive pain in 17; a patient may have had both nociceptive and NeP at the start of the study. To classify pain, some studies [10, 11, 23, 24, 33, 34, 40-42] referred to the taxonomy of SCI pain by Siddall et al. [43]; one study [42] referred to the Bryce-Ragnarsson Pain Classification Scheme [44]. Pain was reported as a characteristic of the population, as an inclusion criterion [10, 11, 23–25, 27, 33, 34, 39–42], an exclusion criterion [38], a limitation for MI [32], an outcome measure [10, 11, 23–25, 27, 33, 34, 37, 39–42], an adverse event [11, 24] or not reported at all [20, 21, 35, 36]. There was high heterogeneity in participants with SCI, who were different in years since lesion, NLI and completeness of injury and type of pain.

Interventions

The imagery capacity of the participants with SCI was tested in three out of 18 studies only, by using KVIQ [25, 32, 35]. Different MI protocols were applied in the studies included. MI interventions were applied with audiotape support [11, 20, 24, 38], under supervision [21, 32, 35], or in combination with brain computer interface (BCI) [36, 37, 40, 41]. Other studies used virtual walking training, which required the participants to imagine performing the movements they were shown, as a standalone intervention [10, 25, 33, 42] or in combination with

transcranial direct current stimulation (tDCS) [23, 34]. One study [39] applied mirror visual feedback in which patients while looking at the reflected image of their non-paralyzed/ unaffected limb in the mirror (occupying the space of their paralyzed/affected or phantom limb) had to perform or imagine the movements of both non-paralyzed/unaffected limb and non-paralyzed/affected or phantom limb.

The duration of sessions varied from 8 min per day to 60 min per day, and total treatment length varied from 1 to 84 days. Follow-up assessments were performed at 1 [21], 2 [35], 3 [23, 32, 33, 38], and twelve months [36].

Outcome measures

The most common measures were pain intensity, measured with VAS and NRS [10, 11, 23-25, 33, 34, 39-42]. Further pain measures location of were the pain [10, 11, 23, 24, 34, 40], pain quality (description of painsuperficial or deep [39], McGill Pain Questionnaire [10, 37], a scale inspired by the McGill Pain Questionnaire [27], Neuropathic Pain Scale (NPS) [42], Neuropathic Pain Symptom Inventory [23]), and the temporal aspects of pain such as the duration of pain relief [10]. The Brief Pain Inventory (BPI) [23, 40], the Basic Pain Data Set [25], the Hospital Anxiety and Depression Scale [25], and the Patient's Global Impression of Change (PGIC) [23] were also reported.

Motor component and activity/disability outcomes were assessed with various tests: gait velocity [38], Performance Oriented Mobility Assessment [38], SCI independence measure [38], muscle strength [20, 35], rate of movement [20], kinematics of upper limb [21, 32, 35], Box and Block test [32, 35], Minnesota Manual Dexterity Test [32, 35], muscle strength [21], and Functional Independent Measure [21].

Neurophysiological measurements of brain activity during MI were done in six quasi-experimental [11, 20, 34, 35, 40, 41] and three observational studies [32, 36, 37]. Two studies used fMRI [11, 20], four EEG [36, 37, 40, 41] and one MEG [35].

Review findings

Table 2 presents a summary of the pain intensity outcomes, the motor function and activity/disability related outcomes.

Pain outcomes

The effects of MI interventions on pain severity are conflicting. Studies using visual illusion combined with MI (Fig 2.1), showed either an improvement after the intervention [10, 23, 33, 34, 39, 42] or no effect [25]. Two studies by Gustin et al. using MI supported by audiotape

		Study design	Experimental studies (F	RCT, quasi-experiment.	al)							
Interface 1		Studies	Cramer et al. [20]	Gustin et al. [24]	Gustin et al. [11]	Kumru et al. [34]	Mateo et al. [35]	Richardson et al. [42] ^a	Sharp et al. [38] ^a	Soler et al. [23] ^a	Vuckovic et al. [40]	Xu et al. [41]
Home in the interval of		JBI evidence level	2.c	2.c	2.c	2.c	2.c	1.c	1.c	1.c	2.c	2.c
quadratical control contro control control control control control control cont		JBI quality assessment score (Total/Maximum)	(6/6)	(6/L)	(<i>6</i> / <i>L</i>)	(8/9)	(6/6)	(10/13)	(9/13)	(9/13)	(6/L)	(6/1)
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	opulation	Total number of SCI individuals (N)	10	15	П	38	9	59	18	39	19	14
NB NB<		Age (mean (SD) [range])	30 (13)	42 (12)	48 (15)	47 (12)	30 (8)	45 (11)	54 (12)	45 (16)	45 (9)	NR
			NR	[26-67]	[26-72]	[25–69]	NR	[22–69]	[26-69]	[21-66]	NR	[18-55]
		Gender (f:m)	NR	0:15	2:9	13:25	2:4	12:47	3:15	9:30	5:14	NR
		Years since lesion (mean (SD),	5 (4.7)	13 (10)	17 (16)	6) 6	1.1 (0.7)	15 (11)	>1	8 (8)	11 (6)	12 (8)
		[range])	NR	[2-32]	[2-46]	[0.3 - 40]	[0.5-2.5]	[0.6-40]	NR	[1-31]	[2-25]	[2-33]
		Neurologic level of injury (traumatic or non-traumatic)	CS - Th10 NR	ThI-Th11 NR	Th1-Th10 NR	C5-Th12 (traumatic and non-traumatic)	C6-C7 NR	Tetraplegia, paraplegia (traumatic)	NR	C3–Th12 NR	ThI-L1 NR	Th2-L1 NR
		Complete/ Incomplete; AIS (A, B, C, D)	A and B	A	V	A-D	A and B	Complete and incomplete	Incomplete	A and B	A and B	A-D
		Acute/Chronic pain	N/A	NR	NR	Chronic	N/A	Chronic	N/A	Chronic	Chronic	Chronic
		At/Below-level neuropathic pain	N/A	Below	Below	At/below	V/N	At/below	N/A	At/below	Below	NR
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	Intervention	Type of intervention (site)	MI (for right foot and tong	MI (for right ankle)	MI (for right ankle)	tDCS + VI with MI	Supervised MI (for upper limb movement)	Virtual walking with MI	MI and overground gait training	tDCS + VI with MI	MI (for hands and feet)	MI (for hands and feet)
14 sestion14 sestion21 sestion21 sestion21 sestion21 sestion1 sesti		Dosage	60 min	8 min	8 min	20 min	45 min	20 min	30 min	20 min	180 trials	180 trials
Table <th< td=""><td></td><td></td><td>14 sessions</td><td>21 sessions</td><td>21 sessions</td><td>10 sessions</td><td>15 sessions</td><td>1 session</td><td>24 sessions</td><td>10 sessions</td><td>1 session</td><td>1 session</td></th<>			14 sessions	21 sessions	21 sessions	10 sessions	15 sessions	1 session	24 sessions	10 sessions	1 session	1 session
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $			7 days	7 days	7 days	14 days	35 days	1 day	56 days	10 days	1 day	1 day
	Comparator	Population	Healthy	SCI without NeP	Healthy	SCI without NeP, Healthy	Healthy	N/A	N/A	N/A	SCI without NeP, Healthy	Healthy
		Intervention	N/A	N/A	N/A	N/A	Supervised MI of geometric fo	rms Virtual wheeling with MI	Overground gait training	tDCS VI Control	N/A	N/A
	Outcome	Baseline	Before	Before	Before	Before	Before	Before	Before	Before	Before	Before
Metr Long termMetr InterventionMetr		During intervention	NR	During	During	NR	NR	NR	NR	NR	NR	NR
		After intervention	After	NR	NR	After	After	After	After	After	After	After
Main outcome ⁶ Moor function and neurophysiologial measurements Pain and neurophysiologial measurements Pain and neurophysiologial measurements <td></td> <td>Long term</td> <td>NR</td> <td>NR</td> <td>NR</td> <td>NR</td> <td>8 weeks post intervention</td> <td>NR</td> <td>NR</td> <td>2, 4 and 12 weeks post intervention</td> <td>NR</td> <td>NR</td>		Long term	NR	NR	NR	NR	8 weeks post intervention	NR	NR	2, 4 and 12 weeks post intervention	NR	NR
Secondary outcome functionNRNRSommosensory functionNRAbsorption in clictsNRAbsorption in clictsNRNAISENRNAISENRNAISENRNAISENRNAISENRNAISENRNAISENRNAISENRNAISENRNAISENRNAISENR <t< td=""><td></td><td>Main outcome^c</td><td>Motor function and neurophysio-logical measurements</td><td>Pain</td><td>Pain and neurophysio- logical measurements</td><td>Pain</td><td>Motor function, activity and neurophysio-logical measureme</td><td>Pain</td><td>Motor function and activity</td><td>Pain</td><td>Pain and neurophysio- logical measurements</td><td>Pain and neurophysio- logical measurements</td></t<>		Main outcome ^c	Motor function and neurophysio-logical measurements	Pain	Pain and neurophysio- logical measurements	Pain	Motor function, activity and neurophysio-logical measureme	Pain	Motor function and activity	Pain	Pain and neurophysio- logical measurements	Pain and neurophysio- logical measurements
Study designObservational studies (case reports, case series)According to the field of		Secondary outcome ^c	NR	NR	NR	Somato-sensory function	NR	Absorption in virtual reality	NR	Anxiety and adverse effects	NR	NR
Study IDGrangeon et al.Grangeon et al.Grande et al. <t< td=""><td></td><td>Study design</td><td>Observational stud</td><td>lies (case reports, case</td><td>series)</td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></t<>		Study design	Observational stud	lies (case reports, case	series)							
IBI evidence level 4d 4d 4c		Study ID	Grangeon et al. [21] ^b	Grangeon et al. [32]	Katayama et al. [33] ^b	Moseley	[10], part a Moseley [10], part b	Onose et al. [36]	Roosink et	al. [26]	Salisbury et al. [37]	Sumitani et al. [39]
JBI quality assessment score (Total/ (68) (78) (68) (7/10) (7/10) (8/10)		JBI evidence level	4.d	4.d	4.d	4.c	4.c	4.c	4.c		4.c	4.c
Population Number of SC1 individuals (N) 1 1 1 1 1 5 4 9 9 25 Age (mean (SD) [range]) 41 23 22 32 (8) 34 (9) 33 (11) 53 (13) 45 (13) NA NA NA NA NA NA 124-45] 124-45] 123-51] 12-5-72] 118-64] Gender (Em) 0:1 0:1 0:1 0:1 0:1 0:1 6:19		JBI quality assessment score (T Maximum)	^r otal/ (6/8)	(7/8)	(6/8)	(1/10)	(7/10)	(8/10)	(8/10)		(8/10)	(6/10)
Age (mean (SD) [range]) 41 23 22 32 (8) 34 (9) 33 (1) 53 (13) 45 (13) NA NA NA NA NA 124-45] 123-51] 125-72] 118-64] Gender (f:m) 0:1 0:1 0:1 0:1 0:4 1:8 2:7 6:19	Population	Number of SCI individuals (N)	1	-	1	5	4	6	6		25	2
N/A N/A [24-45] [23-51] [25-72] [18-64] Gender (i:m) 0:1 0:1 0:1 0:1 0:1 0:5 0:4 1:8 2:7 6:19		Age (mean (SD) [range])	41	23	22	32 (8)	34 (9)	33 (11)	53 (13)		45 (13)	42 and 62
Gender (i:m) 0:1 0:1 0:1 0:5 0:4 1:8 2:7 6:19			N/A	N/A	N/A	[24-45]	[24-45]	[23-51]	[25-72]		[18-64]	N/A
		Gender (f:m)	0:1	0:1	0:1	0:5	0:4	1:8	2:7		6:19	1:1

	Study design	Observational stu	udies (case reports, case :	series)						
	Study ID	Grangeon et al. [21] ^b	Grangeon et al. [32]	Katayama et al. [33] ^b	Moseley [10], part a	Moseley [10], part b	Onose et al. [36]	Roosink et al. [26]	Salisbury et al. [37]	Sumitani et al. [39]
	Years since lesion (mean (SD), [range])	2.6 N/A	0.6 N/A	5 N/A	1 (6) [5–20]	13 (6) [6-20]	NR [0.5–16]	7 (3) NR	median time $= 0.2$	0.6 and 4.8 N/A
	Neurologic level of injury (traumatic or non-traumatic)	C6 NR	C6 (traumatic)	C2 NR	Th12 - L3 NR	L1 - L3 NR	C4 - C7 NR	C3 -C5 to L2 - L3 NR	C, Th and L NR	C and Th NR
	Complete/Incomplete; AIS (A, B, C, D)	¥	A	V	В	В	A - C	A - D	Complete and incomplete	Incomplete
	Acute/Chronic pain	N/A	N/A	Chronic	Chronic	Chronic	N/A	NR	N/A	NR
	At/Below-level neuropathic pain	N/A	N/A	Below	At/below	At/below	N/A	At/below	N/A	NR
Intervention	Type of intervention	MI (for right and left arm)	MI Visual and kinesthetic (for upper limbs)	Virtual visual feedback with MI (while placing the patient on a tilted table)	VI with MI (Virtual walking)	VI with MI (Virtual walking)	MI (in training phase)	Interactive virtual walking with MI (Virtual walking)	BCI MI movement (Cube rotation game, pushing o rolling the cube)	Mirror visual feedback r (Visuomotor imagery)
	Dosage	30 min 10 sessions 14 days	45 min 15 sessions 35 days	2 training periods with 12 weeks washout period First: 10 min	10 min 1 session 1 day	10 min 15 sessions 15 days	30 min 1–2 sessions NR	90 min 2 sessions 1 week between sessions	12 trials of 8s each NR	10 min once a day 4 and 24 weeks
				36 sessions 84 days Second: 10 min 18 sessions 42 days						
Comparator	Intervention	Physical rehabilitation	N/A	Placing the patient on a tilt table	Watching an animated comedy film	N/A	N/A	Static virtual scene during virtual walking with MI	N/A	N/A
Dutcome	Baseline	Before	Before	Before	Before	Before	Before	Before	NR	Before
	During intervention	NR	NR	NR	NR	NR	NR	NR	NR	NR
	After intervention	After	After	After	1 h after	After	After	After	NR	After
	Long term	1 month post intervention	1 and 3 months post intervention	After 4, 8, and 12 weeks	NR	3 months	6 and 12 months post intervention	NR	NR	NR
	Main outcome ^c	Motor function	Motor function	Pain	Pain	Pain	Neurophysio-logical measurements	MI vividness, effort and speed	Neurophysio-logical measurements	Pain
	Secondary outcome ^c	NR	NR	NR	Duration of pain relief	Duration of pain relief	Clinical variables/factors (discomfort/ trouble)	Ongoing pain intensity (pre-post change), perceived interaction with the avatar and virtual environment and adverse effects	Mood, pain, adverse effects	NR
JBI (Th other str and oth designs	e Joanna Briggs Institute idy designs, 1.c–RCT, 2r lower study designs, 2 (3.a–systematic review) Levels of E 1.d—pseudo- 2.c—Quasi-ex of comparat	vidence: Level 1 RCTs); Level 2– sperimental prosl ble cohort studie	-Experimental Design -quasi-experimental de pectively controlled stu .ss, 3.b-systematic revi	s (1.a—Systemai signs (2.a—syste dy, 2.d—pre-test iew of comparat	tic review ematic revi t-post-test ble cohort	of Randomized Con ew of quasi-experim or historic/retrospec and other lower st	trolled Trials (RCTs), 1.b- nental studies, 2.b-system tive control group study); udv designs, 3.c-Cohort		of RCTs and xperimenta nal-analytic roup, 3.d-

JBI (The Joanna Briggs Institute) Levels of Evidence: Level 1—Experimental Designs (1.a—Systematic review of Randomized Controlled Trials (RCTs), 1.b—Systematic review of quasi-experimental other study designs, 1.c—RCT, 1.d—pseudo-RCTs); Level 2—quasi-experimental designs (2.a—systematic review of quasi-experimental studies, 2.b—systematic review of quasi-experimental and other lower study designs, 1.c—RCT, 1.d—pseudo-RCTs); Level 2—quasi-experimental designs (2.a—systematic review of quasi-experimental studies, 2.b—systematic review of quasi-experimental and other lower study designs, 2.c—Quasi-experimental prospectively controlled study, 2.d—pre-test-post-test or historic/retrospective control group study); Level 3—observational-analytic designs (3.a—systematic review of comparable cohort studies, 3.b—systematic review of comparable cohort and other lower study designs, 3.c—Cohort study with control group); Level 4—observational–descriptive studies (4.a—Systematic review of designs, 3.e—Observational study, 3.e—Observational study without a control group); Level 4—observational–descriptive studies (4.a—Systematic review of designs, 5.c—Bench research/single expert opinion) 4.c—case series, 4.d—case study); Level 5—Expert opinion and Bench research (5.a—systematic review of expert opinion; 5.b—expert consensus, 5.c—Bench research/single expert opinion) standard deviation, tDCS transcranial direct current stimulation, VI visual illusion

^aRCT randomized control trials

^bCase study with cross-over design

^cOutcomes of the original studies

Outcome	Study design	Outcome measure	Study ID	2	Population	Intervention group	Before MI (mean (SD) if not stated otherwise)	During MI (mean (SD) if not stated otherwise)	After MI (mean (SD) if not stated otherwise)	<i>P</i> value (within the same intervention group)	<i>P</i> value (between different intervention groups)
PAIN	Experimental studies	NRS (0-10)	Kumru et al. [34]	18	SCI with NeP	tDCS + VI	7.8 (0.9)	NR	4.9 (2.0)	<0.05 (before vs after MI)	N/A
	(NC1, Quasi- experimental)	(01-0)	Richardson et al. [42]	59	SCI with NeP	Virtual walking	NR	NR	– 1.2 (0.3) (Mean change (SEM))	<0.0001 (before vs after Virtual walking with MI)	0.3 (Virtual walking vs Virtual wheeling)
						Virtual wheeling	NR	NR	– 0.5 (0.3) (Mean change (SEM))	0.07 (before vs after Virtual wheeling with MI)	
			Soler et al. [23]	29	SCI with NeP	tDCS + VI	7.5 (1.2)	NR	5.2 (1.5)	<0.05 (before vs after tDCS + VI)	0.008 (tDCS + VI vs VI) 0.004 (tDCS + VI vs control)
						VI	7.2 (1.6)	NR	6.4 (1.6)	<0.05 (before vs after VI)	
						Control (VI without images of human movement)	7.1 (1.5)	NR	6.6 (1.9)	>0.05 (before vs after control)	
		VAS (0–10)	Gustin et al. [24]	2	SCI with NeP	MI of right foot	2.9 (0.7)	5.0 (1.0)	NR	<0.01 (before vs during MI)	N/A
			Gustin et al. [11]	Ξ	SCI with NeP	MI of right foot	3.2 (0.7)	5.2 (0.8)	NR	<0.01 (before vs during MI)	N/A
	Observational studies (case reports, case	NRS (0–10)	Sumitani et al. [39]	6	Participant with SCI #1	Mirror visual feedback	5	NR	Good (pain relief of > 50%)	NR	NR
	series)			_	Participant with SCI #2	Mirror visual feedback	8	NR	Poor (pain relief of < 30%)	NR	
		VAS (0–100)	Katayama et al. [33]	-	SCI with phantom imb pain: left arm	Virtual walking	71 mm	NR	48 mm	<0.05 (before vs after Virtual walking)	NR
					SCI with phantom imb pain: ight arm	Virtual walking	71 mm	NR	53 mm	<0.05 (before vs after Virtual walking)	
			Moseley [10], part a	ŝ	SCI with NeP	Virtual walking	NR	NR	-42 mm [-73 to -11] (mean [95%CI])	NR	NR
						Control (watching animated film)	NR	NR	- 4 mm [- 11 to - 3] (mean [95%CI])	NR	
			Moseley [10], part b	4	SCI with NeP	Replicated case series of Virtual walking	NR	NR	- 53 mm [- 61 to - 45] (mean [95%Cl])	NR	N/A
			Roosink et al. [26]	6	SCI with NeP	Virtual walking	3.3 (3)	NR	3.1 (2.8)	NR	N/A
Motor function and activity related	Experimental studies (RCT, Quasi- experimental)	Maximum tapping rate of tongue (Hz)	Cramer et al. [20]	10	SCI	MI (Practiced task)	1.4 (0.9)	NN av	1.8 (0.2)	<0.0005 (before vs after MI)	NR
		Mucolo atronoth	Connor of al [20]	9	5	MII (Unpracticea task) MI	1.2 (U.4) 0	NR	0.1	 Manual (Delate VS alier IMI) 	N1/A
		Muscle strengtn (Newton)	Cramer et al. [20]	2	20	IM	5	NK	Ð		N/A
		Wrist extension angle during grasping (°)	Mateo et al. [35]	9	SCI	MI	18 (5)	NR	27 (19)	<0.001 (before vs after MI)	N/A
		BBT (number)	Mateo et al. [35]	9	SCI	MI	24 (14)	NR	26 (14)	1.00 (before vs after MI)	N/A
		MMDT (minute)	Mateo et al. [35]	9	SCI	MI	136 (88)	NR	144 (110)	0.53 (before vs after MI)	N/A
		Muscle strength (MMT)	Mateo et al. [35]	9	SCI	MI	5	NR	5	NR	N/A
		Gait velocity (cm/sec)	Sharp et al. [38]	18	SCI	OT + MI	55 (38)	NR	62 (40)	0.005 (before vs after OT + MI)	$\begin{array}{c} 0.27 & (OT + MI \ vs \\ OT) \end{array}$
						OT	41 (32)	NR	56 (51)	0.005 (before vs after OT)	
		POMA (0-28)	Sharp et al. [38]	18	SCI	OT + MI	17 (6)	NR	18 (7)	NR	NR
						OT	16 (9)	NR	18 (7)	NR	
		SCIM (0-100)	Sharp et al. [38]	8	SCI	OT + MI	82 (12)	NR	83 (12)	NR	NR
	Obcomissional attribution	U and trainatour.	Conserve of al	-	5	UT DTAIT MIRT	/6 (12) NB	NK	79 (11) Simifount officer of hoth	NK 0.005 and 0.008 an u.and	div
	Ubservational studies (case reports, case series)	Hand trajectory variability	Grangeon et al. [21]	-	20	LIVIN OF MALLY	NK	NK	Significant effect of both rehabilitation procedures in the horizontal plane	u.u. and u.u.z on x and y axis	XX

 Table 2 Pain and motor function/activity related outcome measurements results.

Outcome	Study design	Outcome measure	Study ID	×	Population	Intervention group	Before MI (mean (SD) if not stated otherwise)	During MI (mean (SD) if not stated otherwise)	After MI (mean (SD) if not stated otherwise)	P value (within the same intervention group)	<i>P</i> value (between different intervention groups)
		FIM (18–126)	Grangeon et al.	-	SCI	PT/MI or MI/PT	49	NR	52	NR	NR
		MT (ms)	[21] Grangeon et al. [32]	-	SCI	MI IL side	2350 (100)	NR	1600 (150)	<0.001	NR
						MI CL side	2300 (300)	NR	1700 (150)	<0.001	
		BBT trained side (number)	Grangeon et al. [32]	-	SCI	MI	25	NR	28	NR	N/A
		MMDT trained side (minute)	Grangeon et al. [32]	-	SCI	IM	3.3	NR	2.1	NR	N/A
BBT Boy Manual 1 reported.	t and Block Test, C Muscle Test, MT M NRS numeric rating	71 confidence interv lovement time testin g scale. 07 overgro	al, <i>CL</i> contralat ig the action of a	eral, J graspi DMA	<i>FIM</i> Function ing the glass	al Independence Meas in the IL space and the mance Oriented Mobil	sure, <i>IL</i> ipsilate e CL space, <i>N</i> n lity Assessment	ral, <i>MI</i> motor number of par <i>PT</i> Physical	imagery, MMDT M ticipants, N/A not app Training BCT Band	innesota Manual Dex olicable, NeP neuropa	terity Test, MMT thic pain, NR not

injury, SCIM Spinal Cord Injury Independence Measure, SD standard deviation, SEM standard error, tDCS transcranial direct current stimulation, VAS visual analog scale, VI visual illusion

[11, 24] showed an increase in pain intensity during the intervention that was maintained for a period of 40 min after the end of the practice (Fig 2.2). In Gustin et al. [24], when patients reported an increase in pain, it was within the same area of the usual ongoing pain. In Gustin et al. [11], when pain increased (9/11 participants), the pain was still located within the usually painful area for six participants, but was spread outside that area in three.

The duration of pain relief was reported only in one study by Moseley et al. [10] as a second outcome measure. Pain relief was lasting longer and the area of pain diminished in size after 15 days of virtual walking training program using MI.

Sumitani et al. [39] categorized the pain descriptions into two main types: superficial pain for "nociceptive pain and temperature sensation" or deep pain for "pain related to pressure sensation and the proprioceptive sense of movement and posture" [39 p. 1039]. They observed a significant decrease in the counts of deep pain linked to visuomotor imagery. In Moseley et al. [10], the pain quality, determined with the McGill Pain Questionnaire, did not change as a result of the intervention. Patients with NeP reported their pain as stabbing, cutting, burning, stinging, and intense [10]. Other studies used the McGill Pain Questionnaire but did not report the quality of pain [27, 37]. Richardson et al. [42] reported that patients with NeP experienced a significant reduction in pain unpleasantness (as measured with the NPS) and a change in certain sensory qualities of that pain ("cold", "deep pain") when compared with the control condition (Fig 2.3). The BPI was also used to assess the intensity and the location of pain prior to the intervention [40]. In Soler et al. [23], pain interference with activities of daily living was assessed using the BPI. They reported the greatest improvement at the end of treatment in the group with tDCS and visual illusion in comparison with the three other groups (tDCS, visual illusion, and placebo) (Fig 2.3). They also reported a significant decrease in anxiety after the last treatment in all intervention groups, as well as pain relief, using the PGIC, after the last day of treatment for all patients [23].

Motor function and activity/disability outcomes

Motor function and activity/disability outcomes were assessed in five studies [20, 21, 32, 35, 38]. Conventional therapy was used in addition to MI in all five. One study [38] assessed lower limb function and four [20, 21, 32, 35] upper limb function. Because of high heterogeneity of the studies, the data for motor function and activity/disability outcomes are presented in narrative form in Table 2.

Cramer et al. [20] showed that 1 week of MI training produced greater gains on maximum physical tapping rate of tongue and right foot, for a practiced than for an

2.1 Comparing effect of MI on pain intensity at baseline and post-treatment

	Ba	selin	е	Post-t	reatm	ent	Mean Difference		Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	\$D	Total	IV, Random, 95% CI		IV, Random, 95% CI
2.1.1 Virtual walking and t	DC S								
Kumru 2013 [34]	7.8	0.9	18	4.9	2	18	2.90 [1.89, 3.91]		
Soler 2010 [23]	7.5	1.2	10	5.2	2	10	2.30 [0.85, 3.75]		
2.1.2 Virtual walking									
Katayama 2015 [33]	7.1	0	1	4.8	0	1	Not estimable		
Moseley 2007, part B [10]	5.6	1	4	1.8	0.9	4	3.80 [2.48, 5.12]		
Roosink 2016 [25]	3.3	3	9	3.1	2.8	9	0.20 [-2.48, 2.88]		
Soler 2010 [23]	7.2	1.6	9	6.4	1.6	9	0.80 [-0.68, 2.28]		
								⊢ -10	-5 0 5 10

2.2 Comparing effect of MI on pain intensity at baseline and during intervention

	Ba	selin	e	During	interver	ntion	Mean Difference		Mean Di	ifference	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Random, 95% CI		IV, Rando	m, 95% Cl	
Gustin 2008 [24]	2.9	0.7	7	5	1	7	-2.10 [-3.00, -1.20]		\rightarrow		
Gustin 2010 [11]	3.2	0.7	11	5.2	0.8	11	-2.00 [-2.63, -1.37]				
								10			
								-10	-0 Fovoure [hoseline]	U D Favours (durin	u interv1
									r avours [basenne]	r avours [uurii	ig interv.j

2.3 Comparing effect of MI to control intervention on pain intensity

	Co	ontro	1	Expe	rimen	ital	Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Random, 95% CI	IV, Random, 95% CI
2.3.1 Virtual walking and t	DCS							
Soler 2010 [23]	6.6	1.9	10	5.2	1.5	10	1.40 [-0.10, 2.90]	
2.3.2 Virtual walking								
Moseley 2007, part A [10]	-0.4	0.6	5	-4.2	1.9	4	3.80 [1.87, 5.73]	
Richardson 2019 (42)	-0.5	1.6	29	-1.2	1.5	30	0.70 [-0.09, 1.49]	+
Soler 2010 [23]	6.6	1.9	10	6.4	1.6	9	0.20 [-1.37, 1.77]	—
								· · · · · ·

Ó 5 10 -5 Favours [control] Favours [experimental]

Fig. 2 Forests plots presenting the results of studies investigating the effect of motor imagery (MI) interventions on pain intensity in individuals with spinal cord injury. 2.1 Five studies were included, comparing the effect of an MI intervention on pain intensity at baseline and post treatment, which used either virtual walking (VW) combined with transcranial direct current stimulation (tDCS) (VW + tDCS) or VW by itself. Three studies showed statistically significant results in favor of treatment by reducing pain and two showed nonsignificant

unpractised task. Sharp et al. [38] showed improvement in gait velocity both in a group using only overground training (OT) and in a group using OT in combination with MI. In their case study, Grangeon et al. [21] reported elbow extensor muscle scores increased by 1 point (maximal score 5) after MI (five sessions a week for 2 consecutive weeks). They also found an increased elbow amplitude associated with a decreased shoulder amplitude from pre-test to follow-up after MI and physical training [21]. Grangeon reduction of pain. 2.2 Two studies, comparing the effect of an MI intervention on pain intensity at baseline and during treatment, were included in this group, which had results showing an increase in pain intensity during MI. 2.3 Three studies were included comparing MI and a comparator intervention in terms of effect on pain intensity, with two in favor of the MI intervention. 95% CI 95% confidence interval, IV inverse variance, MI motor imagery, SD standard deviation, tDCS transcranial direct current stimulation, Total number of participants.

et al. [32] found that the movement time and trajectory smoothness of the upper limb improved following training and those measures remained stable after 3 months. Mateo et al. [35] showed a clinically significant improvement of wrist extension during tenodesis grasping after MI combined with usual rehabilitation, but no other effects on kinematics.

It was not possible to extract information about motor function outcomes after MI intervention separately for individuals with complete and incomplete SCI, or for tetraand paraplegic individuals, because these data were either pooled, not provided at all or there was a big discrepancy in the number of those with complete and incomplete SCI.

Neurophysiological outcomes

Several studies used neuroimaging, such as fMRI (with three-dimensional voxel analysis), to evaluate brain activity during MI intervention in individuals with SCI. It was shown that brain areas involved in movement control undergo reorganization after SCI [20]. Particularly, people with SCI, when compared with healthy individuals, showed the following changes: (1) extended activation volume in the left globus pallidus and posterior putamen-areas of the basal ganglia, which are involved in storage of learned motor sequences and in preparation for motor execution [45] and (2) spatial localization shifts of the primary sensory cortex activation area. Similar brain restructuring was reported in studies by Gustin et al. [11] and Mateo et al. [35], who observed a greater number of voxels activated in the supplementary motor area (involved in both initiation and inhibition of movements [46]), premotor cortex (involved in planning of movement [45]), and cerebellar cortex (involved in motor preparation, and particularly in inhibition of motor commands [45]) of individuals with SCI compared with healthy controls. It was also shown that MI training decreased the threshold for motor system activation after the application of transcranial magnetic stimulation [20]. Analysis of sensorimotor cortex activity using MEG showed that after MI intervention there was a decrease in brainwaves of beta frequencies (13-35 Hz, the range relevant for human brain motor processes) and in event-related desynchronization (ERD) parameters reflecting cortical excitation [35]. Some studies also reported dense neural connections between motor cortex and brain regions involved in pain processing [11, 34]. In contrast to SCI patients without NeP, who had reduced ERD-EEG [35], SCI patients with NeP had an increase in ERD-EEG [40, 41]. This higher EEG activity in SCI patients with NeP was associated with better BCI performance than in those without NeP [36, 37, 40]. Evaluation of electro-dermal response duration (which positively correlates with motor cortex activity) during actual and imagined movement showed that SCI patients can perform MI as accurately as nondisabled people [32].

Adverse events

Eleven studies [10, 11, 20, 23–25, 34, 36, 37, 39, 40] mentioned adverse events but none listed serious adverse events. Two [37, 39] specified that there were no adverse events related to the intervention. An increase in pain was

reported in five studies [11, 23–25, 40] for 20 participants. For 15 participants [11, 24], the increase of pain intensity had already been reported as an outcome measure. The increase in pain was transient and pain returned to its preintervention intensity within 40 min after the intervention [11, 24]. Unpleasant sensations (paresthesia, dysesthesia) were reported by eight participants [24, 40]. Other adverse events included headache (n = 4) [23, 36], fatigue (at least ten participants) [20, 23, 25, 36], difficulty maintaining attention (n = 2) [25], mild transient postural hypotension (n = 1) [36], and distress during virtual walking (n = 1) [10].

Discussion

The objectives of this systematic review were to synthetize therapeutic benefits of MI on pain and motor function recovery in individuals with SCI, as well as to review neurophysiological outcome measures, and to describe the optimal type and dosage of MI intervention.

Although high heterogeneity in studies did not allow us to do meta-analysis and draw the firm conclusions, certain observations were made.

First, most of the findings confirmed the benefits of MI interventions on motor function when combined with physical practice [20, 32, 35, 38]. The results from three RCTs [23, 38, 42] showed positive effects of MI as the sole treatment, on pain or motor function, but also when MI was used as an adjunct to other treatment (e.g., visual illusion, tDCS, or overground training).

Second, multiple repetitions and sessions of 30–60 min were used for motor function improvement [20, 21, 27, 32, 35, 38] and sessions of 8–20 min for pain treatment [10, 11, 23, 24, 33, 34, 39, 42].

Third, pain reduction was observed when comparing the effects within (but not across) the groups before and after MI intervention, whatever the protocol of MI treatment [10, 23, 25, 33, 34, 42]. However, this was not the case in studies comparing pain before and during MI intervention [11, 24]; then an increase in pain intensity was reported. There could be many reasons behind these results. For example, it is possible that MI, as well as the report of actual pain, is a complex cognitive process, which requires competencies and resources of the participant. Therefore, evaluation of actual pain should not be performed during MI [15, 47, 48]. Other causes might be related to the specifics of spinal cord lesion (i.e., NLI, severity, years since injury etc.), MI abilities of the individuals with chronic pain (in the presence of pain MI decreases over time [27]), and/ or type of pain assessment.

Indeed, imagery capacity of the participants with SCI was tested only in three out of 18 studies by using the KVIQ

[25, 32, 35]. In those studies, general MI ability was very variable across the participants, ranging from poor to good, and with greater visual than kinesthetic imagery ability. These results point towards the importance of assessing imagery ability before an MI intervention as it could influence the results. In addition, we examined if the differences in pain outcome results between studies evaluating pain during and after MI intervention could be explained by NLI, the type and subtype of pain, or the pain assessment. The NLI was reported in all studies, except one [38] and varied. The pain type (nociceptive, neuropathic) and subtype (at or below level) were not always reported. Both NLI and pain subtype seem to have unclear impact on an MI intervention's effect on pain. Duration of pain (acute versus chronic) could also play role in MI interventions' effect. For example, a systematic review of different populations with musculoskeletal pain reported a significant MI effect only in those with chronic pain [49]. No information could be found in relation to the years with pain after SCI, but in one study participants with longer duration of SCI experienced a greater reduction in pain intensity from pre- to posttreatment [42]. In addition, there was a lack of consistent definitions of SCI pain categories in the studies included. Clear pain classification for individuals with SCI, as proposed in ISCIP (International Spinal Cord Injury Pain Classification), is considered an important step to correctly assess and treat the pain [2, 50]. Also, pain assessments across the studies were not the same. According to the Initiative on Methods, Measurement, and Pain Assessment in Clinical Trials recommendations about core outcome measures for chronic pain clinical trials [31], various dimensions of pain should be evaluated, such as pain intensity, pain quality, and the temporal components of pain.

In addition, other important outcomes of pain studies include physical and emotional functioning, patient satisfaction with treatment, symptoms, and adverse events [31]. For example, it was shown that anxiety and depression in individuals with SCI and pain [51] might affect MI performance [52–54]. However, in our systematic review we could not evaluate these psychosocial variables, as they were not explored in the studies retrieved.

When considering the effects of MI interventions on motor function, it could be important to test if different MI protocols should be applied for SCI individuals with complete and incomplete injury, tetra- or paraplegic patients. However, no conclusions could be drawn from the studies for the reasons of data heterogeneity or impossibility to extract results separately for individuals with complete and incomplete SCI. Similarly, only a few studies investigated the relationship between the completeness of SCI and pain outcome [23, 42]. They found no association between general pain changes [23] or changes in pain severity [42] and level of SCI (lumbar, thoracic, or cervical) or SCI severity (complete and incomplete). Others studies either did not consider this question or information could not be extracted.

Some studies included in this review examined neurophysiological measures to check brain activity when performing MI. Their data showed significant cortical reorganization after SCI [55, 56], when compared with nondisabled people. The changes were with respect to brain activation volume and patterns both during MI and movement execution. Particularly in people with SCI and NeP, dense neural connections were reported between motor cortex and brain regions involved in pain processing [11, 34]. Interestingly, MI interventions reduced the number of recruited neurons, which could partially explain the motor function recovery and decrease in pain [32]. It was also observed that in contrast to SCI patients without NeP, those with NeP had higher EEG activity and better BCI performance [36, 37, 40]. However, it was not clear if this higher EEG resting state reflected abnormal activity in pain matrix brain circuitry, caused by cortical reorganization, or was a result of antidepressant and antiepileptic medication often taken by persons with SCI with NeP [40, 57].

Conclusion

Based on this systematic review, we cannot give detailed MI intervention guidelines or protocols for pain and motor function recovery in the SCI population. Only general observations can be offered, such as:

- (1) It seems that when performing MI, pain outcome is not influenced by the level or severity of SCI.
- (2) Shorter MI sessions were applied for pain reduction (average time of about 15 min based on reported range of 8–20 min) than for motor/functional improvement (average time of about 45 min based on reported range of 30–60 min). Therefore, to create guidelines more studies are needed with similar protocols with respect to population, intervention, and outcomes.

We think that the design of an MI intervention should also take into account the following factors, which were addressed by Milton et al. [58]: (1) complexity of the motor task and challenging environment are important to get better results, because conditions closer to a real world environment engage the motor system in an optimal way; (2) best performance requires attention to the assigned task as well as the ability to filter irrelevant information, which might be impaired in patients with nervous system diseases. In addition, before performing MI, it is crucial to test the imagery capacity of the participants [15] as neurological deficits may affect it [8].

Future research

There should be more studies comparing MI pain and motor function outcomes between individuals with complete and incomplete SCI. The effects of MI interventions on pain and its stability over time remains questionable. Therefore, clinical trials evaluating MI as standalone and/or adjunct therapy for NeP in SCI patients are warranted to develop appropriate guidelines for MI treatment.

Data availability

The datasets generated during the current study are available from the corresponding author upon request.

Funding The study was in part supported by a grant from the University of Applied Sciences and Arts Western Switzerland//HES-SO (scientific commission of health) to Emmanuelle Opsommer (73642/S-RAD17–04).

Author contributions EOP and NKO were responsible for designing the review protocol, writing the protocol and report, conducting the search, and screening potentially eligible studies. EOP, OCH, and NKO were responsible for critically appraising studies, extracting and analyzing data, and interpreting results, updating reference lists, creating Summary of findings' tables, and writing the final report.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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