



Associations between socioemotional alterations, quality of life, and social functioning in multiple sclerosis: A scoping review

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Abstract

The main aim of the present scoping review is to systematically review the available studies that investigated the associations between socioemotional alterations (i.e., social cognition impairments/alexithymia/difficulties in emotion regulation) and both reduced QoL and social functioning in patients with Multiple Sclerosis (MS). The articles were selected from the PubMed, PsycINFO, and Scopus databases. The main exclusion criteria were qualitative studies, articles that did not use validated instruments, and studies that did not investigate the association between socioemotional skills and QoL/social functioning in MS. Of the eight studies fulfilling the inclusion criteria, six found significant associations between social cognitive and emotion regulation abilities and QoL/social functioning in patients with MS, while two found no significant relationships particularly between the performance on social cognition tasks and QoL measures. Overall, the majority of findings seem to highlight that socioemotional alterations contribute to impaired QoL and social functioning in MS. However, given the still limited evidence, future studies are needed to replicate and confirm the available results, paying attention to two principal aspects: the use of standardized and ecological tasks for the assessment of social cognition skills and the recruitment of samples involving patients with different types of MS.

Keywords Multiple sclerosis · Social cognition · Emotion regulation · Alexithymia · Quality of life · Social functioning

Introduction

Multiple sclerosis (MS) is a chronic inflammatory and neurodegenerative disease of the central nervous system, which affects approximately 2.5 million individuals all over the world (Files et al., 2015). It is usually diagnosed between the age of 20 and 40 and its prognosis is often unpredictable. Depending on the location and extent of lesions, patients may experience different disease-related stressful conditions, such as motor weakness, sensory deficit, impaired balance, and urinary disturbance.

In addition to physical symptoms, patients with MS often report cognitive impairments (Calabrese, 2006) and high levels of psychological distress (Feinstein, 2004; Siegert &

Abernethy, 2005). Particularly, psychiatric comorbidities can occur in up to 95% of patients with MS. Depression is the most common condition with a prevalence of around 50%, followed by anxiety, which ranges from 14 to 41% (Paparigopoulos et al., 2010).

Similarly, cognitive decline has been reported in up to 65% of patients with MS (Benedict et al., 2017; Messinis et al., 2018). Main signs of cognitive impairment usually include deficits in information processing speed, verbal and visuospatial memory, verbal fluency, and executive functions (Planche et al., 2015; Prakash et al., 2008).

More recently, researchers have started to focus their attention also on the investigation of social and interpersonal aspects of cognition (i.e., social cognition) in patients with MS. Social cognition has been defined as ‘the ability to construct representations of the relation between oneself and others and to use those representations flexibly to guide social behavior’ (p. 231, Adolphs, 2001). Examples of social cognition abilities include both the capacity to represent other people’s intentions and beliefs (i.e., Theory of Mind, ToM; Premack & Woodruff, 1978;

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Leslie, 1987), and the ability to share and recognize the emotions of others (Lieberman, 2007).

Deficits in social cognition have been reported in a number of clinical conditions, such as autism spectrum disorder, schizophrenia, neurodegenerative disorders and chronic pain conditions (e.g., Senju, 2013; Bora et al., 2015; Di Tella et al., 2015; Green et al., 2015), compromising the patient's ability to implement appropriate social interactions. Alterations in social cognition skills have also been linked to difficulties in both the ability to identify and describe one's own feelings (i.e., 'alexithymia'; Taylor et al., 1997), and the ability to correctly regulate one's own emotions (e.g., Le Berre, 2019; Di Tella et al., 2020).

All those competencies together are fundamental to obtain adequate social support from other people and to cope with chronic disabling conditions such as MS (Chalah & Ayache, 2017a). Conversely, if these skills are compromised, people may find themselves struggling to interact effectively in interpersonal contexts, with negative consequences on Quality of Life (QoL) and social functioning (Krause et al., 2013; Schwartz & Frohner, 2005). While QoL has been defined by the World Health Organization as an 'individuals' perception of their position in life in the context of the culture and value systems in which they live, and in relation to their goals, expectations, standards and concerns', social functioning usually refers to the ability to fulfill the individual's role within his/her different environments, such as work and social environments (e.g., Bosc, 2000).

Furthermore, social cognition deficits and alexithymia have been shown to be significantly associated with anxiety/depressive symptoms (Luminet, 2009; Richards et al., 2002; Washburn et al., 2016), which in turn can contribute to the high prevalence of affective disorders observed in patients with MS.

Previous systematic reviews have been carried out to address the magnitude and extension of social cognition impairments and alexithymia in MS studies. The majority of those reviews have focused on the measurement and neuroanatomical correlates of social cognition and alexithymia (e.g., Bora et al., 2016; Cotter et al., 2016; Chalah & Ayache, 2017b; Labbe et al., 2018), highlighting the presence of alterations in ToM (especially when assessed through visual tasks), recognition of negative facial emotional expressions (anger and fear), and ability to adequately identify and describe one's own feelings.

However, the possible effects of these cognitive and emotional deficits on QoL and social functioning of patients with MS have not previously been integrated.

Therefore, the main aim of this scoping review is to systematically review the available studies that investigated the association between socioemotional alterations (i.e., social cognition impairments/alexithymia/difficulties in emotion

regulation) and both reduced QoL and social functioning in patients with MS.

Methods

The aim of a scoping review is to identify and summarize key concepts in a particular research area. In contrast to a systematic review, the quality of evidence is not evaluated within a scoping review. Instead, broader topics and research questions are addressed with the purpose of identifying research gaps and making recommendations for future research. The present scoping review was conducted following the Preferred Reporting Items for Systematic reviews and Meta-Analyses extension for Scoping Reviews (PRISMA-ScR) checklist (Tricco et al., 2018). The protocol was not pre-registered.

Search Strategy

A comprehensive, systematic literature search was conducted between November 2020 and January 2021 in PubMed, PsycINFO, and Scopus databases. The following keywords combinations were used to identify target published articles: (social cognition OR emotion recognition OR theory of mind OR empathy OR alexithymia OR emotion regulation) AND (quality of life OR social functioning) AND multiple sclerosis.

The last run was conducted on the 14th January 2021. With the use of this search string, 1.270 titles were initially identified from 1942 up until 2021 (see Fig. 1 for the flow diagram of article selection).

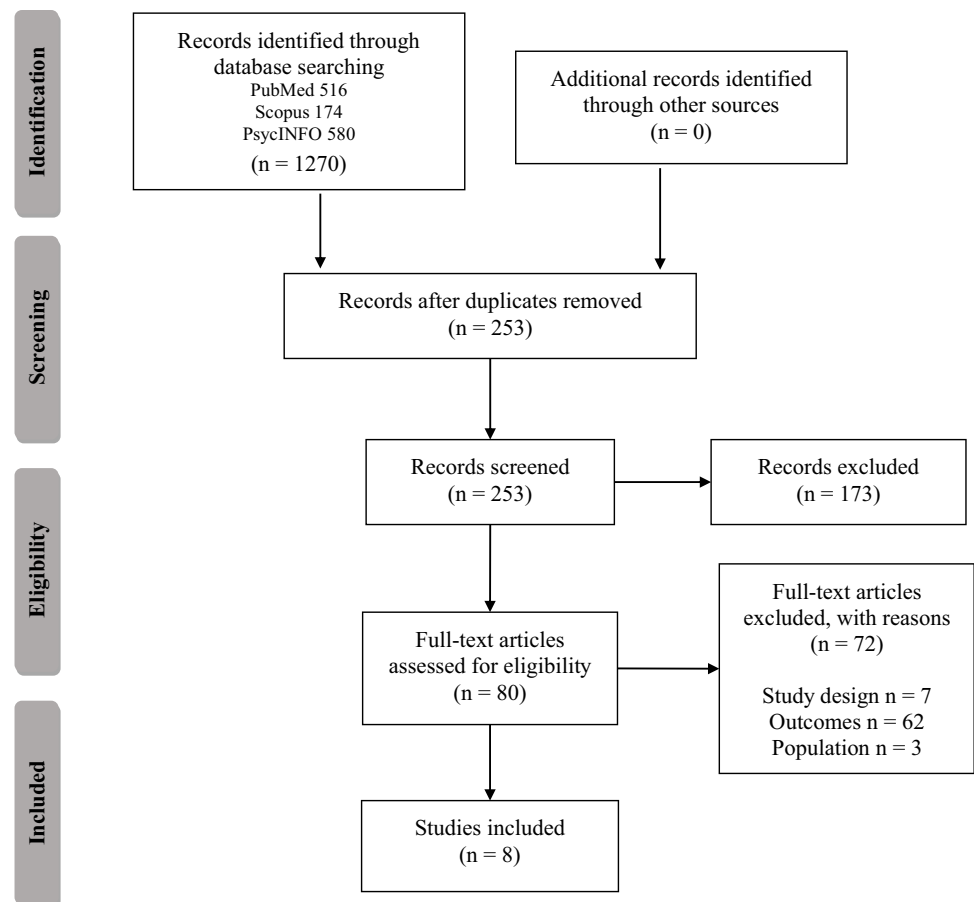
Eligibility Criteria

The present review aimed to identify peer-reviewed academic articles that investigated the relationship between social cognition/alexithymia/emotion regulation and QoL/social functioning in patients with MS. Only academic peer-reviewed scientific papers that were published in English were eligible for inclusion in this review.

Exclusion criteria were as follows:

- 1) Articles that exclusively assessed social cognition (i.e., ToM, empathy, and emotion recognition) impairments or alexithymia or emotion regulation, without providing further insights on the association between those deficits and QoL/social functioning in patients with MS.
- 2) Studies that used ad hoc constructed instruments or qualitative methods.
- 3) Articles that recruited and assessed patients with MS under 18 years of age or patients with MS together with patients with other neurological conditions.

Fig. 1 Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) flow diagram for the study selection



- 4) Articles published but not peer reviewed or under review at the time the search was carried out.
- 5) Textbooks and book chapters were also excluded, because many textbooks and book chapters do not provide new insights, as original research papers do, and not all of them are peer reviewed.

Study Selection

Study selection was carried out by two authors (MDT and LC), who first screened the titles and abstracts of all the retrieved articles and then examined the full text of all remaining publications. Disagreements on the inclusion or exclusion of publications were discussed until agreement was reached. Reference lists of relevant articles were also checked, in order to obtain additional important publications that were not included yet.

Data Extraction

The characteristics of all included studies were extracted by one author (MDT). Data items that were extracted from each included study were: author and year of publication, participants, methods, and main findings.

Results

Study Selection

After duplicates were removed, a total of 253 records were screened. Of these articles, 245 titles have been excluded based on title, abstract, and full text screening (Fig. 1). Most of these excluded articles did not assess social cognition skills (ToM, emotion recognition, and empathy), alexithymia or difficulties with emotion regulation in patients with MS ($n = 173$) or did not evaluate those constructs with regard to QoL/social functioning ($n = 62$) or did not use a research design that allowed to investigate those associations ($n = 7$). The remaining articles ($n = 3$) enrolled mixed samples of patients, including people with MS and other neurological conditions. The study selection process resulted in 8 studies that fulfilled the inclusion criteria and were included in the present review.

Characteristics of Included Studies

A summary of the main characteristics and results of the 8 studies included in the present review is provided in Table 1. The selected articles were published from 2009 to 2020,

Table 1 Summary of the selected studies concerning the multivariate relationship between multiple sclerosis, socioemotional skills, and quality of life/social functioning

Authors (year)	Participants	Measures	General results	Correlations with QoL/SF
<i>Social cognition</i>				
Ciampi et al. (2018)	23 PP; 20 SP MS patients (28F; mean age: 57.2, DD: 15.7) No control group	<i>Social cognition</i> : mini-SEA <i>Cognition</i> : BICAMS, Stroop test <i>Mood</i> : BDI-II <i>QoL</i> : MSIS-29	Poor cognitive performance at the BICAMS Low levels of depressive symptoms Low levels of QoL	No significant correlations between QoL and social cognition tasks
Grothe et al. (2020)	50 NP MS patients (29F; mean age: 39.4, DD: NP) No control group	<i>Social cognition</i> : emotion recognition test, RMET <i>Cognition</i> : BRB-N <i>Mood</i> : BDI-II <i>QoL</i> : MusiQoL	Cognitive performance on average at the BRB-N Low levels of depressive symptoms Low levels on "Relationships with friends" dimension of MusiQoL	No significant correlations between QoL and social cognition scores
Isernia et al. (2019)	26 RR; 8 SP; 8 PP MS patients (24 F; mean age: 52.4, DD: 21.2) 26 HCs	<i>Social cognition</i> : RMET, Strange Stories, Faux pas, MASC <i>Cognition</i> : MOCA, BRB-N, PARADISE 24 <i>Mood</i> : BDI-II, STAI-Y1 <i>QoL</i> : MSQOL-54	Poor performance on Strange Stories and Faux Pas tests High levels of depressive symptoms Poor cognitive performance at the MOCA Lower scores at the physical domain of QoL than HCs	Significant main effect of Strange Stories (contrasting emotion passage) on both the physical and mental health domains of QoL
Phillips et al. (2011)	27 RR; 2 PP; 3 SP MS patients (28 F; mean age: 44.0, DD: 7.9) 33 HCs	<i>Social cognition</i> : static Ekman facial expressions and brief dynamic emotion perception video clips; BFRT, as control task <i>Cognition</i> : FAS letter fluency task, memory task (from SEFCI), and SART for inhibition <i>Mood</i> : HADS <i>QoL</i> : WHOQoL-BREF	Deficits in facial emotion recognition No difference at the control task Poor cognitive performance (FAS and SART) Lower scores in all domains of QoL than to HCs	Positive and significant correlations between both measures of emotion perception and the psychological and social aspects of QoL. Those associations were found to be still significant, after controlling for severity and duration of disease, age, and cognition
van der Hiele et al. (2020)	278 RR MS patients (216 F; median age: 43.0, DD: 5.0) 128 HCs	<i>Social cognition</i> : Empathy Quotient <i>Cognition</i> : PASAT, SDMT, RAVLT, BVM-T-R, TMT, D-KEFS subtests <i>Mood and Personality</i> : HADS, NEO-FFI <i>Occupation</i> : WPAL, WRFQ-2.0, WRFQ-2.0, MSWDQ-23	No difference on empathy scores Poor cognitive performance Higher scores on depressive symptoms and neuroticism; lower scores on extraversion, openness, and conscientiousness than HCs Poor occupational functioning	Empathy did not differ between employed and unemployed individuals with MS Positive and significant correlations between empathy scores and occupational functioning (work scheduling and output demands and cognitive/psychological work barriers)
<i>Emotion regulation</i>				
Phillips et al. (2009)	61 RR; 17 PP/SP; 8 NP MS patients (63 F; mean age: 44.8, DD: 11.8) No control group	<i>Emotion regulation</i> : ERQ <i>Cognition</i> : CFQ <i>QoL</i> : WHOQoL-BREF	Low scores on emotion regulation Moderate scores on the attention and QoL measures	Significant predictor role of ERQ reappraisal scores for psychological and environmental dimensions of QoL. For social QoL, the ERQ-reappraisal approached significance

Table 1 (continued)

Authors (year)	Participants	Measures	General results	Correlations with QoL/SF
Phillips et al. (2014)	27 RR; 2 PP; 3 SP MS patients (216 F; mean age: 44.0, DD: 7.9) 32 HCs	<i>Emotion regulation</i> : DERS <i>Cognition</i> : FAS, SART <i>Mood</i> : HADS <i>QoL</i> : WHOQoL-BREF	Difficulties in emotion regulation abilities Poor cognitive performance Higher scores on depressive symptoms than HCs Lower levels of QoL than HCs	Negative and significant correlations between DERS scores and psychosocial dimensions of QoL
Schirda et al. (2015)	85 RR; 7 SP; 3 PP MS patients (79 F; mean age: 43.7, DD: 10.4) No control group	<i>Emotion regulation</i> : DERS <i>Mood and Mindfulness</i> : BDI-II, FFMQ <i>QoL</i> : WHOQoL-BREF, SWLS	Difficulties in emotion regulation abilities Moderate levels of depressive symptoms High scores on QoL dimensions	Negative and significant correlations between DERS total score and composite QoL score

Age and disease duration are expressed in years

BDI Beck Depression Inventory, *BICAMS* Brief International Cognitive Assessment for MS, *BFR* Benton Facial Recognition Test, *BRB-N* Brief Repeatable Battery of Neuropsychological Tests, *BVMT-R* Brief Visuospatial Memory Test-Revised, *CFQ* Cognitive Failures Questionnaire, *DERS* Difficulties in Emotion Regulation Scale, *DD* disease duration, *D-KEFS* Delis-Kaplan Executive Function System, *ERQ* Emotion Regulation Questionnaire, *FFMQ* Five Facet Mindfulness Questionnaire, *HADS* Hospital Anxiety and Depression Scale, *HCS* healthy controls, *MASC* Movie for the Assessment of Social Cognition, *mini-SEA* Social Cognition and Emotional Assessment, *MS* multiple sclerosis, *MSIS-29* Multiple Sclerosis Impact Scale 29, *MSQOL-54* Multiple Sclerosis Quality of Life 54, *MSWDQ-23* Multiple Sclerosis Work Difficulties Questionnaire, *MusiQoL* Multiple Sclerosis International Questionnaire of Quality of Life, *NP* not provided, *PASAT* Paced Auditory Serial Attention Test, *PP* primary progressive, *QoL* Quality of life, *RAVLT* Rey Auditory-Verbal Learning Test, *RR* relapsing remitting, *RMET* Reading the Mind from the Eyes, *SART* Sustained Attention to Response Task, *SEFCT* Screening Examination for Cognitive Impairment, *SD* standard deviation, *SDMT* Symbol Digit Modalities Test, *SF* Social Functioning, *SP* secondary progressive, *STAI-YI* State-Trait Anxiety Inventory – Y1, *SWLS* Satisfaction with Life Scale, *TMT* Trail Making Test, *WAI* Work Ability Index, *WPAI* Work Productivity and Activity Impairment Questionnaire, *WRFQ-2.0* Work Role Functioning Questionnaire 2.0

with half of the studies being carried out in the last four years ($n=4$).

Most of articles reported about the type of MS and disease severity. Research samples typically include a combination of people with relapsing and progressive types of MS and involve patients with various levels of disability.

Based on the different aspects the selected articles focused on, we differentiated between studies that investigated the relationship between social cognition skills (i.e., ToM, emotion recognition, and empathy) and QoL/social functioning ($n=5$), and studies that examined the association between alexithymia/emotion regulation and QoL/social functioning ($n=3$). The two article categories are presented below, separately.

Social Cognition, Quality of Life, and Social Functioning in Multiple Sclerosis

Among the studies that investigated the presence of social cognition deficits in patients with MS, five articles included in their assessment also the analysis of QoL and social functioning aspects. Those studies obtained, at least in part, distinct results. Indeed, two studies did not find significant correlations between lowered social cognition and QoL scores (Ciampi et al., 2018; Grothe et al., 2020), while the remaining studies found significant associations between reduced social cognition performance and worse QoL (Isernia et al., 2019; Phillips et al., 2011) or between altered social cognition and reduced social functioning (van der Hiele et al., 2020).

Particularly, the study of Ciampi et al. (2018) employed the mini-Social Cognition and Emotional Assessment (mini-SEA) (Bertoux et al., 2012, 2014) to assess different aspects of social cognition in a group of patients with progressive types of MS ($n=43$). The mini-SEA consists of two different tests, namely a shortened version of the Faux-Pas test and the Face Emotion Recognition test. The Faux Pas test includes both short passages, in which a character inadvertently hurts or offends another, and control stories, in which no faux pas is committed. The individual is required to identify if the faux pas has been made, why the main character did it (cognitive ToM) and how the victim of the faux pas must have felt (affective ToM). The Face Emotion Recognition test consists of 35 pictures, representing the six basic emotions (i.e., happiness, sadness, anger, surprise, fear, and disgust) and neutral expressions. The individual is asked to select the correct option for each of the presented photographs.

Among the different objectives the study pursued, the association between social cognition skills and QoL is of particular interest to the present review. QoL aspects have been assessed by means of the Multiple Sclerosis Impact Scale (MSIS-29; Hobart et al., 2001), which is a 29-item

self-report questionnaire consisting of 20 items associated with the physical dimension of QoL and 9 items associated with the psychological component of QoL. Results of correlation analyses showed no significant association either between mini-SEA scores and QoL scores, or between faux pas and face emotion recognition subtest scores and MSIS-29 scores.

Similarly, the study of Grothe et al. (2020) investigated the associations between social cognition abilities, clinical variables, and QoL in a cohort of patients with MS ($n=50$). For the assessment of social cognition skills, the authors administered the following two tests: (1) a facial morphing task to assess emotion recognition, which consists of 48 morphing sequences (from neutral to the target expression), presented in random order (Lischke et al., 2012). The participants have been required to press a button when they had recognized one of the possible four emotions (i.e., happy, angry, fearful, sad); (2) the Reading the Mind in the Eyes (RMET; Baron-Cohen et al., 2001) to evaluate affective ToM, which requires the individual to choose the correct mental state among four alternatives that are provided below each of the 36 images the test is made of. With regard to QoL evaluation, the Multiple Sclerosis International Questionnaire of Quality of Life (MusiQoL; Simeoni et al., 2008) was administered. It allows the investigation of nine dimensions of QoL, which are specifically related to MS.

In order to examine the relationships between social cognition measures and QoL dimensions, correlation analyses were performed. Results showed the presence of non-significant associations between social cognition tasks and MusiQoL total or subscale scores.

Different results were found by the studies of Isernia et al. (2019) and Phillips et al. (2011). Particularly, Isernia et al. (2019) aimed to investigate the relationship between ToM, clinical variables, and neuropsychological deficits in a sample of patients with both relapsing and progressive types of MS ($n=42$), compared to a group of healthy controls (HCs; $n=26$). For the assessment of cognitive and affective ToM processes, the authors employed the following 4 tasks: (1) the RMET (Baron-Cohen et al., 2001); (2) the Strange Stories test (Happé, 1994), which includes both mental and physical passages that require the participant to infer characters' mental states or to make global inferences that go beyond what was explicitly mentioned in the text, respectively. For mental stories both total score and single-story scores have been considered; (3) Faux pas test (Baron-Cohen et al., 1999), of which the authors selected 4 faux pas and 4 matched control stories; (4) the Movie for the Assessment of Social Cognition (MASC) (Dziobek et al., 2006), which consists of 15-min video clips representing an ecological situation in which four friends meet to have dinner together. Video is interrupted several times to ask participants multiple-choice questions on emotions, thoughts, and intentions

of the different characters involved in the story. In addition to ToM tasks, the Multiple Sclerosis Quality of Life 54 questionnaire was administered for the assessment of QoL dimensions (Vickrey et al., 1995).

With particular regard to social cognition and QoL, the authors tested whether a poor ToM performance was related to low levels of QoL. General Linear Model was performed with QoL scores as within-subject factor and ToM scores as between-subject factor. Results of GLM showed a main effect of contrasting emotion passage of the Strange Stories on both the physical and mental components of QoL. No other significant association between ToM tasks and QoL measure was detected. Those findings suggest a significant role that affective ToM seems to play in different areas of QoL.

Yet the study of Phillips et al. (2011) examined the ability to identify emotional and non-emotional information and whether difficulties in emotion perception were related to QoL in a cohort of patients with both relapsing and progressive types of MS ($n = 32$) compared to a group of HCs ($n = 33$). To assess emotion perception, the authors administered the following four tests: (1) static images of facial emotion, taken from the standard set of Ekman facial expressions, were used to assess the ability to correctly identify expressions of the six basic emotions (Young et al., 2002); (2) the Benton facial identity recognition test (Benton et al., 1983) was included, as a control task, to examine non-emotional aspects of facial perception; (3) dynamic emotion perception video task (Slessor et al., 2007), in which participants were asked to select the word that best describes the feelings of the main character in 16 videos; (4) dynamic age and gender perception video task was employed as a control task for the dynamic emotion perception task. With regard to QoL, the authors administered the World Health Organization quality of life questionnaire (WHOQoL-BREF; Skevington et al., 2004) to investigate both scores of patients with MS on four domains of QoL (i.e., Physical, Psychological, Social, and Environmental) and their relationship with difficulties in emotion perception. Results showed that people with MS rated their QoL as poorer than HCs in all domains of functioning. Furthermore, correlation analyses revealed that both measures of emotion perception were strongly related to psychological and social domains of QoL, but not to the physical domain. Perceiving emotions from videos was also associated with the environmental domain of the WHOQoL-BREF. Correlations between emotion perception and QoL were found to be still significant after controlling for disease severity and duration, age, and cognition, suggesting a specific pattern of associations between difficulties in emotion perception and poor social and psychological QoL.

Finally, the study of van der Hiele et al. (2020) investigated the possible associations between empathy and

cognitive, psychological, and occupational functioning in a group of patients with relapsing–remitting MS ($n = 278$) compared to HCs ($n = 128$). For the assessment of empathy the authors administered to all participants the Empathy Quotient (Baron-Cohen & Wheelwright, 2004), which is a 60 item self-report questionnaire with total score ranging from 0 to 80 (higher scores indicate greater empathic abilities), while for the evaluation of occupational functioning various measures have been employed to investigate different work-related aspects (e.g., self-reported influence of MS symptoms on productivity while at work; work role functioning; current work ability as compared to lifetime best; psychological/cognitive, physical, and external work difficulties). With particular regard to the association between empathy and occupational functioning, no significant differences were detected between employed and unemployed individuals with MS on empathy scores. However, results of correlation analyses showed the presence of significant associations between higher empathy scores and better occupational functioning, in terms of work scheduling and output demands, and less cognitive/psychological work barriers in patients with MS. These findings suggest that reduced empathy could be negatively associated with worse occupational functioning, with negative consequences on the individual's ability to perform complex activities of daily life.

Alexithymia, Emotion Regulation, Social Functioning, and Quality of Life in Multiple Sclerosis

Similar to social cognition skills, only three articles included in their assessment the analysis of QoL dimensions in relation to emotion regulation abilities (Phillips et al., 2009, 2014; Schirda et al., 2015). Conversely, to the best of our knowledge, no study has investigated the relationship between alexithymia and either social functioning or QoL in patients with MS yet. Likewise, we did not retrieve any article that examined the association between emotion regulation abilities and social functioning.

The three available articles yielded similar results in terms of significant associations that were detected between difficulties in emotion regulation and reduced QoL. Particularly, the study of Phillips et al. (2009) mainly aimed to investigate the association between emotion regulation strategies and QoL in a group of people with both relapsing and progressive types of MS ($n = 86$). For the assessment of emotion regulation, the authors employed the Emotion Regulation Questionnaire (ERQ; Gross & John, 2003), which allows to evaluate two different types of emotion regulation strategies, that is reappraisal and expressive suppression, while for the assessment of QoL they administered the WHOQoL-BREF (Skevington et al., 2004). In order to examine the relationships between QoL dimensions and emotion regulation abilities, a series of regression analyses were performed. Results

showed that ERQ-reappraisal significantly predicted poor psychological QoL and environmental QoL, while for social QoL, ERQ-reappraisal approached significance ($p=0.06$). Conversely, expressive suppression was not found to be a significant predictor of any of QoL dimensions. These findings seem to highlight that failure to use adequate emotion regulation strategies may contribute to reduced QoL in patients with MS.

Similar findings were obtained by the study of Phillips et al. (2014), which analyzed the associations between emotion regulation difficulties and other factors including QoL in a group of patients with both relapsing and progressive types of MS ($n=31$) compared to HCs ($n=31$). Emotion regulation difficulties were assessed by means of the Difficulties in Emotion Regulation Scale (DERS; Gratz & Roemer, 2004), which is a self-report questionnaire designed to assess multiple aspects of emotion regulation, while the WHOQoL-BREF (Skevington et al., 2004) was employed for the assessment of QoL-related dimensions.

Results of comparisons between patients with MS and HCs revealed that the former reported greater difficulties in emotion regulation and poorer QoL compared to HCs. With regard to the association between emotion regulation and QoL, the authors found that higher scores at the DERS total were significantly correlated with reduced social, psychological, and environmental dimensions of QoL in the MS group only, suggesting that these patients may experience greater difficulties in emotion regulation than HCs, which in turn predict poorer QoL.

Finally, the study of Schirda et al. (2015) examined the relationship between trait mindfulness, QoL, and difficulties in emotion regulation in a group of people with relapsing and progressive types of MS ($n=95$). Similar to the above-mentioned studies, the authors employed the DERS (Gratz & Roemer, 2004) for the assessment of difficulties in emotion regulation, and the WHOQoL-BREF (Skevington et al., 2004) and the Satisfaction with Life Scale (Diener et al., 1985) for the evaluation of QoL-related aspects. With regard to QoL and difficulties in emotion regulation, results of correlation analyses showed that the QoL composite score was significantly and negatively correlated with the DERS total score in people with MS. These findings suggest that MS patients with difficulties in the regulation of emotions experience less QoL.

Discussion

Studies assessing the association between socioemotional alterations and both QoL and social functioning in people with MS are still limited. Indeed, the study selection process only resulted in 8 studies that fulfilled the inclusion criteria and were included in the present review.

The majority of those studies seem to show a significant association between altered socioemotional abilities and QoL/social functioning (Isernia et al., 2019; Phillips et al., 2009, 2011, 2014; Schirda et al., 2015; van der Hiele et al., 2020). Alterations in those domains may have detrimental effects on the individuals' well-being, with negative consequences on QoL and social functioning (particularly, psychological and social dimensions of QoL and work-related aspects). Although the available evidence in MS is still limited, previous studies carried out in other clinical populations seem to support this pattern of results, showing that both social cognition deficits and difficulties with emotion regulation contribute to impaired QoL and social functioning in patients with different medical conditions (e.g., Cooper et al., 2015; Couture et al., 2006; Yogarajah & Mula, 2019).

Despite this positive evidence, not all the results are consistent and other studies found no significant relationships especially between social cognition and QoL dimensions (Ciampi et al., 2018; Grothe et al., 2020).

A possible explanation for those contrasting findings may rely on the heterogeneous instruments that have been employed by the different studies. In fact, whereas for the assessment of emotion regulation abilities and QoL most authors have administered similar self-report questionnaires (e.g., the DERS and the WHOQoL-BREF, respectively), for the evaluation of social cognition skills, various measures have been adopted and this may have led to inconsistent findings. The use of different tasks (e.g., task type and complexity) can also make it difficult to compare results across the studies. In order to overcome this issue, standardized and ecological tasks, which are able to reflect how individuals behave in an enriched social context, should be employed. In this view, dynamic assessment tools, such as videotaped tasks, have been proved to have better accuracy than traditional static tests in assessing social cognition components (Dziobek et al., 2006). For instance, the only study among the ones we included in the present scoping review (Phillips et al., 2011), which used a dynamic emotion perception video task for the assessment of facial emotion recognition, was able to detect a significant association between this measure and psychological, social, and environmental domains of QoL. However, future studies employing instruments, such as the Amsterdam Dynamic Facial Expression Set (Van Der Schalk et al., 2011), the Montréal Pain and Affective Face Clips (Simon et al., 2008) and the Awareness of Social Inference Test (McDonald et al., 2006) for the assessment of social cognition components, are necessary to prove the validity of videotaped tasks to predict QoL and social functioning in people with MS more accurately.

Furthermore, although the evaluation of emotion regulation abilities has been based on the use of similar measures, these instruments are exclusively self-report. The limitations of self-report instruments are well known and

may lead to underestimation of patients' difficulties due to self-reflection impairments typically observed in many neurological disorders. Indeed, explicit self-report measures require the respondents to be aware of their reduced ability to identify and describe feelings (e.g., Parling et al., 2010). Performance-based instruments or structured interviews, less dependent on the patient's awareness, should be employed in addition to traditional self-report measures.

Another possible explanation for the mixed findings the present review showed could be ascribed to the heterogeneous cohorts of people with MS that the different studies recruited. Indeed, patients with different disease subtypes, distinct levels of physical disability, and various stages of illness may be more or less prone to social cognitive deficits, which could have, in turn, a different impact on QoL and social functioning dimensions.

Finally, it is still a matter of debate if social cognition impairments can be considered a result of the cognitive deficits frequently observed in people with MS or if they may represent a primary sign of the disease. Nevertheless, it is clear that the combination of multiple cognitive, social, and affective symptoms may have a leverage effect on the individual's psychological well-being and day-to-day functioning. Indeed, the available evidence seems to show that worse QoL is negatively correlated with greater physical, cognitive and emotional disabilities, and with longer disease duration and progressive clinical course (e.g., Labiano-Fontcuberta et al., 2015; Laing et al., 2015).

The results of the present scoping should also be interpreted with caution due to some limitations of the studies we included. Firstly, most of them adopted a cross-sectional design, which does not allow establishing any causal direction between socioemotional alterations and reduced QoL/social functioning. Secondly, some of those studies did not recruit also a group of HCs to be used as a comparison for MS patients.

Despite those limitations, the present scoping review represents, to the best of our knowledge, the first contribution summarizing the available evidence on the complex relation between altered socioemotional skills and reduced QoL/social functioning in patients with MS, suggesting the presence of significant associations between those constructs in MS population.

Further evidence is, however, necessary to replicate and confirm the available results. A deeper understanding of the alterations in social cognitive and emotional skills may allow researchers and clinicians to have a more comprehensive framework of the psychological symptoms associated with MS. In this way, it would be possible to plan a better medical management with tailored treatments specific for each individual's needs.

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Declarations

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