Patient education for people with multiple sclerosis-associated fatigue: A systematic review

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Abstract

Background

Multiple Sclerosis (MS) is an inflammatory and neurodegenerative disease often causing decreased quality of life, social withdrawal and unemployment. Studies examining the effect of pharmacological interventions demonstrated only minor effects, whereas non-pharmacological interventions as e.g. patient education programs have shown promising results.

Objective

We aim to systematically review the literature to determine the effect of patient education programs on fatigue in MS.

Methods

We conducted a comprehensive search in PubMed for randomized controlled trials (RCTs) that evaluated patient education programs for MS-related fatigue. Interventions evaluating physical exercise and/or pharmacological treatments were not included. Meta-analyses were performed using the generic inverse variance method.

Results

The search identified 856 citations. After full-text screening we identified ten trials that met the inclusion criteria. Data of 1021 participants were analyzed. Meta-analyses showed significant positive effects on fatigue severity (weighted mean difference -0.43; 95% CI -0.74 to -0.11) and fatigue impact (-0.48; -0.82 to -0.15), but not for depression (-0.35 (95% CI -0.75 to 0.05; p = 0.08). Essentially, we categorized patient education programs into two types: firstly, interventions with a focus on cognitive-behavioral therapy (CBT) and secondly, interventions that teach patients ways of managing daily fatigue. CBT-based approaches seem to generate better results in reducing patient-reported fatigue severity. Analysing CBT studies only, the
pooled weighted mean difference for fatigue severity was \(-0.60\) (95% CI; \(-1.08\) to \(-0.11\)) compared to non-CBT approaches \((-0.20;\) 95% CI; \(-0.60\) to \(-0.19\)). Furthermore, interventions employing an individual approach seem to reduce fatigue more effectively than group-based approaches (pooled weighted mean difference for fatigue severity in face-to-face studies was \(-0.80\) (95% CI; \(-1.13\) to \(-0.47\)) compared to group-based studies with \(-0.17\) (95% CI; \(-0.39\) to \(0.05\)). Longest follow-up data were available for 12 months post-intervention.

**Conclusion**

Overall, included studies demonstrated that educational programs and especially CBT-based approaches have a positive effect on reducing fatigue. Since fatigue is thought to be a multidimensional symptom, it should be treated with a multidimensional approach targeting patients' behavior as well as their emotional and mental attitude towards fatigue. However, the clinical relevance of the treatment effects i.e. the relevance for patients' daily functioning remains unclear and long-term effects, i.e. sustainability of effects beyond 6 months, warrants further work. This review has been registered in the PROSPERO international prospective register of systematic reviews data base (Registration number: CRD42014014224).

**Introduction**

Multiple Sclerosis (MS) is a poorly understood disease of the central nervous system that mostly starts in young adulthood [1]. Disease progression is unpredictable and patients may experience a wide range of symptoms. Amongst limb weakness, paraesthesia, visual impairment and other symptoms patients regularly report fatigue. Fatigue is one of the most common symptoms that 75–95% of MS-patients struggle with at least once in their life [2]. Current definitions include a subjective lack of physical and/or mental energy that is perceived by the individual or the caregiver to interfere with usual or desired activity, an overwhelming sense of tiredness, lack of energy or feelings of exhaustion and a feeling of physical tiredness and lack of energy distinct from sadness or weakness [3]. Fatigue is considered to be the most disabling symptom by many patients, more important than mobility restrictions or pain [4]. Fatigue is also a common reason for unemployment [5] and often a cause for social withdrawal [6]. Fatigue may affect patients very differently and causes are yet poorly understood [3].

According to current beliefs, there are primary and secondary mechanisms that lead to fatigue [7]. While primary fatigue is a direct result of physical changes within the body through the disease itself, secondary fatigue is due to other primary factors that can cause and worsen fatigue. Among the primary mechanisms are axonal loss, reorganization and increased brain recruitment as well as immunological and neuroendocrine factors. Factors of secondary fatigue include sleeping problems, depression, stress, side effects of pharmacological therapies and reduced physical activity [8; 9]. Moreover, it has been postulated that fatigue is perpetuated by certain cognitions and beliefs [10].

Currently, the only established way of measuring fatigue is by use of self-reported scales since fatigue is subjective and multidimensional symptom. Most common are scales that measure fatigue severity i.e. the intensity and characteristics of experienced fatigue and/or fatigue impact which focuses on activities that patients can and cannot do due to fatigue. These two measurements do not always correspond because perceived intensities may not have the same impact on different patients' life [11].
Currently there are no convincing pharmacological treatments available for MS-related fatigue [12]. However, there have been a number of non-pharmacological treatment approaches. For example, physical exercise seems to have positive results on fatigue severity [13].

A current systematic review and meta-analysis has suggested promising results of various patient education programs teaching patients fatigue management techniques [14]. The review compared the effects of medication, physical exercise and patient education programs for MS-related fatigue and concluded that physical exercise and education were more effective than pharmacological therapies and should therefore be preferred in fatigue therapy management. The review however only gives a broad overview on different interventions.

As there are various approaches on educating patients and teaching them different skills to better manage and cope with fatigue in daily life, a more focussed review was considered necessary. For this review, we aimed to include two different types of programs: Firstly, programs based on cognitive-behaviour therapy and secondly, programs teaching patients’ general strategies of fatigue management. We aimed to describe the complex interventions in detail, using a recently proposed framework and to analyse the quality of the evidence using the Cochrane risk of bias tool [15] and GRADE methodology [16]. Furthermore, we recorded length and duration of each program as well as settings and implementation strategies. In summary we aimed to determine the best strategy for patient education programs to achieve best results in reducing fatigue.

The objective of the current review is to summarize the results of RCTs assessing the effects of CBT-based patient education compared to other patient education programs that teach ways of managing fatigue in daily life for the reduction of MS-related fatigue in any setting.

**Methods**

This review is based on systematic literature searches in PubMed and the Cochrane library. As we were only looking for RCTs with at least 30 participants, we decided to perform a sensitive search in PubMed as this has been shown to be highly sensitive [17]. Additionally, we searched the Cochrane library and checked reference lists of included studies and other systematic reviews to increase sensitivity. The latest search was conducted on the 3rd January, 2016. We aimed for a maximum sensitive search and therefore only used the search terms “multiple sclerosis” and “fatigue”. The explicit search strategy can be found in the appendix.

**Inclusion criteria**

Only studies that evaluated patient education programs for MS-related fatigue using randomized controlled and crossover designs were included. As education program we defined any intervention where patients receive information or education about strategies to deal with fatigue. Further inclusion criteria were fatigue as a primary outcome measure and a minimum effective sample size of 30 participants. Additionally, we only included studies that assessed fatigue at least at two time points. Studies that included patients with other neurodegenerative/ neuroinflammatory diseases than MS were included if the majority of included patients were MS patients. No limitations concerning publication date, disease course of participants, or language were made.

Eligibility for studies was assessed independently by two reviewers. Discrepancies were solved by discussion afterwards. Reviewers were not blinded to study authors. Data extraction was performed according to a standardized data extraction form. Data for the following items was extracted: (1) sample size, (2) demographic data of participants, (3) type and length of intervention (number of weeks, number of hours per week), (4) type of control group, (5)
Risk of bias assessment

We used the Cochrane Risk of Bias tool to assess study quality. This tool evaluates selection bias, performance bias, detection bias, attrition bias, and reporting bias. Due to the nature of the included studies, it is difficult to blind participants to group allocation since patient education programs require participants to play an active part in the sessions [18]. In combination with all outcome measures being entirely self-reported, this was regarded as a potentially high risk of bias.

Summary measures

All studies were assessed by two independent reviewers. As all outcomes were continuous outcomes, we calculated standardised mean differences (SMD) between groups and 95% confidence interval when sufficient data were available. We contacted authors of studies in case of insufficient information on data at baseline and follow-up. For meta-analyses, we used the generic inverse variance approach using RevMan software [19]. Due to marked clinical heterogeneity, we used random effects models to provide pooled estimated effects.

We additionally performed subgroup analyses by calculating pooled mean differences for trials that worked with a CBT approach and individual vs. group interventions. We define the term ‘individual intervention’ as the session being based on one therapist and one patient in direct contact (face-to-face, via telephone or online).

Risk of bias across studies

We used the specific evidence grading system developed by the GRADE [16] group to assess study quality.

Results

Study selection (see Fig 1)

856 publications were identified by the search. After deleting 78 duplicates, 778 studies remained to be screened by abstract which ultimately led to 30 publications for full-text screening. After reading full-text articles, 20 publications were eliminated for different reasons, namely trials using drugs, sports interventions, magnetic fields, complementary medicine and others. 10 publications that met all criteria were included. For one study we thankfully received additional data from the authors which enabled us to integrate the RCT into our analyses.

Study characteristics (see Table 1)

All included studies were randomized controlled trials published in English, two of which used a crossover design. According to the information provided in the respective publications, group and face-to-face sessions took place in community centres or hotel conference rooms. No interventions were performed in a hospital setting. Interventions took four to ten weeks and follow-up data were available from ten weeks to 12 months. Control groups contained usual care, “placebo” interventions and alternative interventions e.g., relaxation training and physical exercise. Trials were performed in the US (n = 4), Belgium (n = 1), Australia (n = 1), the UK (n = 2), Switzerland (n = 1) and New Zealand (n = 1).
Participants

The number of study participants ranged from 30 to 190. Overall, this review contains data from 1021 participants from 10 trials. In all studies, predominantly women participated which
corresponds to the male: female ratio of patients in MS patients. Mean age of participants ranged from 41.1 to 56 years. Relapsing-remitting MS (RRMS) was the predominantly reported disease course in all studies. Mean years since disease onset ranged from 5.5 to 21 years.

**Interventions (see Fig 2)**

Cognitive-behavioral therapy was the most commonly used approach. Four of ten approaches were based on CBT. Two studies used an energy conservation program. Three studies used...

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**Table 1. Overview of types of interventions, sample sizes, dates, countries and methods of delivery.**

| Country          | Date               | Sample Size | Intervention                                      | Method of delivery                      |
|------------------|--------------------|-------------|--------------------------------------------------|-----------------------------------------|
| Finlayson et al. 2011 [20] | USA 11/2007-04/2009 | 190         | Managing fatigue program                         | Group teleconference                   |
| Ghahari et al. 2010 [21] | Australia 05/2007-03/2008 | 95          | Managing fatigue program                         | Online + online personal contract with facilitators |
| Grossman et al. 2010 [22] | Switzerland n.r. | 150         | Mindfulness-based intervention                   | Group sessions                         |
| Hugos et al. 2009 [23] | USA 03-08/2007     | 30          | Fatigue self-management program                  | Group setting                          |
| Kos et. Al. 2007 [9]   | Belgium n.r. 51    | Multidisciplinary fatigue management program     | Group setting                          |
| Mathiowetz et al. 2005 [24] | USA 2002-3         | 169         | Managing fatigue program                         | Group setting                          |
| Moss-Morris et al. 2012 [25] | UK n.r. 40        | Cognitive-behavioral therapy                     | Online + telephone contact with facilitators |
| Thomas et al. 2013 [26] | UK 11/2007-04/2009 | 164         | Cognitive-behavioral therapy + energy effectiveness training | Group sessions                       |
| Van Kessel et al. 2008 [27] | New Zealand 07/2004-08/2005 | 72          | Cognitive-behavioral therapy                      | Personal contact + telephone sessions  |
| Mohr et al. 2003 [28] | USA n.r. 60        | Cognitive-behavioral therapy                     | Personal contact sessions               |

n.r. = not reported

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**Fig 2. Overview of study length, type of intervention and follow-up.**

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multidisciplinary fatigue self-management programs that informed about different techniques of reducing fatigue e.g. relaxation, cooling, energy saving but also teaching the benefit of physical exercise. One study employed the concept of mindfulness.

Different ways of program delivery were employed. Mainly, two different formats are distinguishable: group programs and individual programs. Group sessions were the most commonly used format. Further, there were different ways of contact between therapist and participant, e.g. personal contact, online and telephone contact. Two interventions used online sessions; one with online personal contact to the facilitators, one combined online with telephone support sessions.

Another difference is the length of intervention in weeks (and weekly hours) as well as length of follow-up. The shortest intervention was 4 weeks and the longest was 16 weeks. Fig 2 gives an overview of the types of program, length of the program and follow-up.

Outcomes

Interventions used different fatigue measures. The most commonly administered outcome measures were the Fatigue Impact Scale [29] and the Modified Fatigue Impact Scale [30], each used in four trials. Further outcome measures were the Fatigue Severity Scale [31] and the Fatigue Scale [32]. Furthermore, depression was assessed in 6 studies and health-related quality of life in 3 studies. Also, some studies assessed anxiety as secondary outcome measurements. All outcomes measures were taken at baseline and directly post-intervention. 6 studies reported follow up data for fatigue impact and 3 studies for fatigue severity (between three and six months).

Risk of bias in individual studies (Fig 3)

We did not detect selection bias in most studies. Due to the nature of patient education interventions, staff and patient blinding was not realized as participants and facilitators engage with each other, but this was considered as low risk of bias. To control for therapist time and attention, van Kessel and colleagues used relaxation training [27]. Outcome assessment was considered as blinded in all studies although all outcomes were self-reported. We assessed outcome data as incomplete if more than 10% of participants’ data was not missing. This was the case in one study. To assess selective reporting we searched for published study protocols. If those were not found, we judged the risk as ‘unclear’ (this applied to 6 studies). Furthermore, we assessed study quality with the GRADEpro tool [16].

Synthesis of results

Six studies measured fatigue severity and five measured fatigue impact. Depression was measured by seven studies; while data for quality of life were only available for two studies, so we decided against a quantitative synthesis. Not all studies provided all data necessary to be included in the meta-analysis. Fatigue severity as well as depression data were available from six studies; fatigue impact data were available from four studies. Among further endpoints were anxiety, QUALYs, and personal wellbeing. These were only measured by single studies and will therefore not be discussed further in this review.

Pooled weighted mean difference for fatigue severity (Fig 4) with a pooled sample of n = 509 in six studies was -0.43 (95% CI -0.74 to -0.11; p = 0.008, high quality evidence). CBT-based approaches resulted in more pronounced treatment effects.

For fatigue impact, the pooled weighted mean difference was -0.48 (95% CI -0.82 to -0.15; p = 0.005, high quality evidence) in four studies with 314 participants (Fig 5).
Depression data were available for six studies with 365 participants showing no significant effect directly after the intervention with a pooled mean difference of -0.35 (95% CI -0.75 to 0.05; p = 0.08, high quality evidence; Fig 6).

We detected clinical heterogeneity between studies e.g. in terms of interventions, study populations and follow-up periods. Also, most analyses showed marked statistical heterogeneity. Therefore, random effects models were used for meta-analyses.

Analyses indicated a larger effect of CBT-based approaches than other fatigue management programs for both fatigue severity and depression (Figs 4 & 6). For fatigue impact, we decided against calculating subgroup analyses as there were not enough reported data.

**Fig 3. Risk of bias in individual studies.** (+) low risk of bias; (?) unclear risk of bias; (-) high risk of bias.

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Furthermore, we analysed the studies that use a face-to-face approach. For fatigue severity and for depression, pooled effect sizes indicated larger effects of individual approaches compared to the group settings (Figs 7 & 8). Again, we decided against analysing subgroup data for fatigue impact due to a lack of data.

Discussion

Fatigue has been identified as a major disabling factor in patients with MS. Accordingly, a number of different pharmacological and non-pharmacological interventions have been developed. This systematic literature review has identified 10 RCTs on educational interventions for MS-related fatigue. Based on the findings of this review, educational interventions seem effective in reducing patient-reported fatigue in the short-term (up to 6 months). There is also some indication that fatigue levels may be reduced over a longer period of time. Follow-up study data from two studies showed sustained reduced fatigue levels after one year [33; 34]. However, more research is warranted.

As usually the case with educational interventions, all included interventions were multi-modal or “complex” interventions comprising different components [35] as e.g. provision of information, interaction with peers, practice of acquired techniques and homework. Also, interventions varied markedly, e.g. in mode of delivery and/or duration. This poses a number of challenges for the synthesis of these interventions. Still, we decided to perform meta-analyses as interventions seemed sufficiently homogeneous in terms of the provided information, setting and the common goal of reducing fatigue. Furthermore, three studies based on the
same fatigue management program: Mathiowetz and colleagues tested the “Managing Fatigue” program for the first time while Ghahari et al. and Finlayson et al. used modifications of the same program.

In 2014, Asano & Finlayson published a systematic review comparing the effectiveness of sports interventions, pharmaceutical treatments and educational programs concerning their benefit on MS-related fatigue [14]. The review showed that, of the three intervention types, educational programs had the largest effect in terms of reducing fatigue. In addition to this finding, our review indicates that interventions based on cognitive behavioural therapy seem to be more effective in reducing fatigue compared to other education programs, although this must be taken with caution considering that analyses are based on a small number of studies. This finding is consistent with the assumption that fatigue is, at least partly, caused and perpetuated by negative thoughts and corresponding behaviour [10]. Furthermore, we found some evidence that face-to-face approaches may be superior to group approaches.

Fig 6. Depression (CBT vs. non-CBT approaches).
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Fig 7. Fatigue severity (subgroups: individual vs. group setting).
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The nature of the complex interventions used does not allow a direct comparison of effects of different approaches. Nevertheless, it seems plausible that effectiveness might be increased by combining different methods of intervention provision e.g. online with telephone support sessions or the combination of telephone and face-to-face sessions. Research on mental health conditions suggests, for example, that internet-based interventions tend to be more effective when combined with personal support. Such support is typically provided via e-mail or telephone, and some studies suggest that even minimally trained technicians can be as effective as trained clinicians in their support [36]. Fully automatized support, such as mobile text messages, also appear to be effective across many health conditions and, when offered adjunctively, could boost the effects of internet interventions [37]. Efficient support can also be provided only “on demand” for those who request [38]. Further investigation is needed to examine how different intervention and support elements can be combined to optimize the effects of complex interventions targeting fatigue.

Our findings also suggest that individual approaches may be superior in reducing fatigue severity compared to a group based approach which may be due to closer interaction with the participant and more possibilities of targeting participant-specific problems. On the other hand, group approaches may be easier to implement in a clinical setting as more patients may work with one facilitator at the same time. Furthermore, patients all work together which provides better opportunities for vicarious learning, peer support and modelling. All of this can increase self-efficacy and support behaviour changes. However, no conclusive data on that particular subject are available. Furthermore, the length of the intervention (measured in weeks) and/or the number of sessions might have an impact on their ability to reduce fatigue. The shortest program lasted 4 weeks with 2 hours of program each week without demonstrating a benefit on participants’ fatigue levels [9]. This may be due to shorter practising time since all programs aim to integrate new behavioral strategies into daily life.

The marked reduction of fatigue levels shown by an online CBT intervention by Moss-Morris et al. [25] suggests further exploration of online CBT interventions based on patients’ individual needs and preferences. Evidence from systematic reviews suggests that computer-tailored and web-based interventions that are tailored to suit individual patient characteristics tend to be more effective than non-tailored or “generic” programs. Such evidence mirrors earlier research, which showed that tailored printed health behaviour interventions tend to
outperform their non-tailored counterparts [39]. Further research is needed to clarify which forms of tailoring are most effective (e.g., personalization, content-matching, provision of feedback) and by which mechanisms tailoring effects unfold (e.g., enhanced cognitive processing of tailored interventions).

A limitation of the studies included in our review concerns the limited duration of follow-up periods, which lasted from 10 to 52 weeks. To gain insight as to whether participants are able to maintain reduced fatigue levels or if there is a need for 'booster sessions' after programs as suggested by some authors, further research with long term outcome measurement is needed.

Concerning limitations of our study, firstly, we only searched the Pubmed and Cochrane databank. Although a single search in Pubmed has shown to be highly sensitive [17], we may still have missed data with this approach. Secondly, our approach of comparing CBT studies to non-CBT studies may be difficult since it is possible to apply principles of CBT without doing pure CBT and some of the non-CBT studies may actually apply CBT techniques as well. So, this may confound the results of our study.

Since MS-related fatigue is reported to be varying individually between patients, all patients should receive the combination of therapy best suited for their individual requirements. Online interventions may be a good compromise between achieving the best possible patient-centred care and actual feasibility of a program in a rehabilitation setting.

This review supports the hypothesis that education programs may be effective in reducing fatigue in patients with MS. Here, CBT-based interventions may be the most effective way to teach patients ways of managing fatigue.

Supporting information

S1 File. Search strategy for PubMed.
(TIF)

S2 File. Finlayson initially requested teleconf data (Cochrane review, May 25 2014).
(DOCX)

S3 File. PRISMA checklist.
(DOC)

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