Depression and multiple sclerosis: Review of a lethal combination

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Abstract—Depression is the most frequent psychiatric disorder in multiple sclerosis (MS) patients. The etiology of depression is multifactorial and likely associated with psychosocial stress, focal demyelinating lesions, and immune dysfunction. Proper diagnosis and severity assessment are critical prior to initiation of therapy. Patients with suicidal ideation should be referred for immediate psychiatric consultation and be closely monitored. While more therapeutic trials for depression in MS are needed, MS patients have been shown to respond to current antidepressant medications and psychotherapy. Unfortunately, patients with MS and major depression or suicidal thoughts are often underassessed and therefore not diagnosed. Unlike other aspects of MS, depression is treatable. Early intervention in depression can prevent declines in quality of life and even death from suicide. This article reviews the unique features, assessment, and treatment of depression in MS. MS care providers should vigilantly assess depression and suicide risk in their patients.

Key words: assessment, depression, morbidity, mortality, multiple sclerosis, outcome, prevention, review, suicide, treatment.

INTRODUCTION

Multiple sclerosis (MS) is the most common progressive neurological disease of young adults and currently affects 350,000 persons in the United States [1–2]. Typically, the course of MS starts with random exacerbations and then slowly becomes progressive. Affective disturbances in MS were first noted by Charcot. In 1874 he observed that the intellectual and emotional faculties of patients with MS were “blunted in their totality” [3]. Depression, in particular, is frequently seen in patients with MS but has only recently received attention in the literature. This article highlights the unique features, assessment, and treatment of depression in MS.

Abbreviations: AHCPR = Agency for Health Care Policy and Research; BDI = Beck Depression Inventory; BDI-FS = BDI-Fast Screen; CBT = cognitive-behavioral therapy; DSM = Diagnostic and Statistical Manual of Mental Disorders; DSM-IV = DSM, Fourth Edition; ECT = electroconvulsive therapy; FAMS = Functional Assessment of Multiple Sclerosis; IFN = interferon; IL = interleukin; MRI = magnetic resonance imaging; MS = multiple sclerosis; POMS = Profile of Mood States; QOL = quality of life; SEG = supportive-expressive group (therapy); SIT = stress inoculation training; SSRI = selective serotonin reuptake inhibitor; STAR*D = sequenced treatment alternatives to relieve depression; TMAP = Texas Medication Algorithm Project; TNF = tumor necrosis factor; VA = Department of Veterans Affairs.

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Depression is the most frequent psychiatric diagnosis in patients with MS [4]. The lifetime risk for depression in patients with MS ranges from 40 to 60 percent [5]. Interestingly, age and sex-adjusted prevalence rates were twice as high in patients with MS compared with patients with other chronic diseases in a Canadian population-based study [6]. Patients with MS also have more severe depressive symptoms compared with patients with other chronic neurological diseases [7–9]. One study found that only about one-third of patients with MS and major depression or suicidal thoughts received treatment by their healthcare providers [10]. Both healthcare providers and patients need further education about the identification of at-risk patients and the initiation of therapy.

A number of reasons may explain the association of MS with depression [11], including—

1. The psychosocial effects of MS disability.
2. The direct effect of lesions on brain structures that are involved in regulating and maintaining mood state.
3. The untoward effects of interferon (IFN)-β for treating MS, which may be associated with mood changes.
4. Immune dysfunction.

**PSYCHOSOCIAL EFFECTS OF MULTIPLE SCLEROSIS**

Depression in MS may be a consequence of the multiple challenges associated with managing a chronic illness. In addition to neurological deficits, MS is frequently associated with losses in vocational status, social roles, sense of control, and participation abilities [12]. The nature of MS is unpredictable and potentially unrelenting. Perceptions regarding the uncertainty in disease, intrusiveness in daily activities, and lack of hope have been associated with depression [13]. Life stress and coping abilities may also mediate psychosocial outcomes [14].

**Physical and Cognitive Impairment**

Global physical impairment has been associated with higher levels of depression in some studies [15–17] but not others [11,18]. At first glance, the relationship between impairment and depression appears to be mixed. However, a clearer relationship between depression and impairment appears when studies use more focused measures of physical impairment. For example, Minden et al. found no link between depression and disease severity at a global level; however, depressive episodes were likely to occur within 1 month of a steroid-treated exacerbation of MS [18]. Similarly, another study found increased prevalence of depression during times of MS exacerbations and increased physical impairment [13].

In addition to temporal variation, certain types of physical disability may be differentially related to depression. For example, a recent study of veterans found that perceived mobility and bladder impairment were not associated with increased risk of depression but reports of perceived bowel impairment and “at least occasional” falls were associated with elevated risk of depression [19]. A possible explanation for these findings comes from Devins et al. who found that the relationship between physical disability and depression is indirect: disability affects psychosocial outcome to the degree that impairment is intrusive and personal control is threatened [20].

Physical and cognitive impairments are differentially related to functional outcomes, which highlights the need for separate consideration of these areas [21]. Persons with high levels of cognitive impairment are less likely to work outside the home, more likely to require assistance with activities of daily living, and more likely to have limited social support [22–23]. Some studies have indicated that patients with MS-related cognitive impairment report higher levels of depression than patients without cognitive impairment [24–25]. Other studies do not support this relationship [26–27]. Some evidence exists that the specific type of cognitive impairment may be differentially related to depression. For example, Kenealy et al. found that MS patients with impaired autobiographical memory were less likely to be depressed than those with intact autobiographical memory [28]. The link between cognitive impairment and depression is further complicated by possible bidirectional influence. Not only may cognitive impairment precipitate depression, but depression in a patient with MS may also result in reduced attention and working memory capacity [29]. In sum, depression and cognition are clearly related independent of physical disability, but more research is required for a better understanding of this link.

**Vocational Changes**

In the general population, increased risk of depression is associated with unemployment, disability, “homemaker” status, and living at or near the poverty level [30]. A recent population-based study found that persons with disabilities were 5.0 times more likely than persons without disabilities to lose their jobs [31]. The rates of depression were higher among persons with disabilities than persons without disabilities; unemployment status explained nearly 30 percent
of the elevated depression found in the group with disabilities. The authors argue that the effects of unemployment and disability on depression are independent and additive.

In the MS literature, the links between unemployment and depression are mixed. Given that a majority of patients with MS lose their jobs and about one-third experience a decrease in standard of living [32–33], vocational and financial losses may mediate the relationship between MS and depression. Unemployment among patients with MS is associated with a lower quality of life (QOL) [34]. Williams et al. reported that unemployment was the strongest predictor of a major depressive episode in persons with MS and that the odds of depression were 3.2 times higher among those who were unemployed [19]. This contrasts with a large community-based sample of patients with MS in which unemployment was not associated with depression [17]. Further research will clarify the relationship between vocation and depression in patients with MS.

Social Changes

Studies have generally found that lower levels of perceived social support are associated with depression in MS [16,25]. A significant portion of patients with MS report qualitative changes in social networks and personal relationships as a result of their disease [16,25,35], including the loss of professional colleagues, diminished contact with social groups, and loss of social independence [36]. Concurrently, many patients with MS become increasingly reliant on care providers and core family members, which often increases caregiver burden [34]. Marital status appears to remain relatively constant after MS diagnosis [32], and many married patients with MS report that they obtain most support from their spouse [37]. However, patients with the highest levels of MS disability perceive less overall support, and those with longer duration of illness perceive less affective and affirmation support than those with shorter duration of illness [37]. A number of studies have found that women and unmarried persons with MS are at particular risk for diminished social support during their illness [36,38]. Moreover, unsupportive relationships are significantly and independently correlated with depression and a lower sense of purpose [39]. Future studies of social support should address the influence of both supportive and unsupportive behaviors.

Coping

Most research on coping in MS has relied heavily on Lazarus and Folkman’s model that describes coping as “constantly changing cognitive and behavioral efforts to manage specific external and/or internal demands that are appraised as taxing or exceeding the resources of the person” [40]. This model (and subsequent modifications) suggests that most coping strategies available to an individual experiencing a stressor fall into two global yet distinct categories: emotion-focused coping and problem-focused coping. Emotion-focused coping most often represents the individual’s reactive efforts to reduce distress caused by the stressor (e.g., avoidance, wishful thinking). Problem-focused coping represents the individual’s active efforts to change a stressful situation through modifications of the environment, his- or herself, or the actual stressor (e.g., information gathering, goal setting).

Several studies have identified a link between emotion-focused coping and poor adjustment in MS; depression was the most common outcome reported. In cross-sectional surveys, emotion-focused coping has been related to lower self-esteem [41], global distress [42–44], and depression [13,15,45–46]. Similar results were found in studies that examined the relationship between coping styles and depression over time [47–48]. One common interpretation of these findings is that emotion-focused coping is ineffective and leads to poorer adjustment. However, others have argued that in a progressive degenerative disease with often limited opportunities for actively reducing disability or symptoms, emotion-focused coping may be useful and be a substantial portion of a patient’s available coping efforts. Furthermore, seemingly passive strategies such as acceptance may have important benefits [12,49].

The relationship between problem-focused coping and depression in MS is less well established. Problem-focused coping has been linked to higher self-esteem [41], global distress [42], and depression [15,45], but other studies have failed to find a correlation [13,43–44]. Longitudinal studies have sometimes found associations between problem-focused coping and depression. This link, however, is often observed only at one of several time points and not when emotion-focused coping is examined simultaneously [47–48]. Despite limited data, problem-focused coping is viewed as particularly promising because it has been associated with well-being in general population samples [40] and because problem-focused strategies are the foundation of many psychosocial interventions. Certain problem-focused coping strategies are also argued to be more helpful than others or only appropriate in particular circumstances [12]. These important nuances may be lost in studies that measure coping with global scales.
BRAIN LESIONS AND MOOD

Generally, changes in mood have been associated with neurological illnesses with a subcortical component. Since MS affects myelin and the integrity of nerve conduction, mood changes may be a direct neurological consequence of the disease. Several studies have looked at the association between brain lesions and mood changes. Di Legge et al. reported findings from a group of patients with clinically isolated syndromes [50]. They monitored patients for relapse and conversion to MS and found an increased rate of depression in patients who were developing MS. They also reported an association between lesions in the right temporal region and depression severity.

Pujol et al. investigated the relationship between depressive symptoms in MS and lesions in the frontal and temporal lobes [51]. Using a standardized magnetic resonance imaging (MRI) protocol, they studied white matter lesions in tracts connecting these two lobes of the brain. Lesions in the suprainsular white matter (arcuate fasciculus) were significantly associated with depressive symptoms.

Mohr et al. reported a relationship between brain lesion volume and depression in patients with MS and major depression [52]. Patients were enrolled in one of three 16-week treatment programs. They were studied at the end of treatment and 6 months later. At the end of treatment, Beck Depression Inventory (BDI) scores were positively associated with right temporal periventricular lesion volume and left temporal gray-white junction lesion volume. At 6-month follow-up, BDI scores were positively associated with total lesion volume, lesion volume in multiple discrete areas, and neuropsychological functioning.

Bakshi et al. used MRI to study the relationship between lesion location, brain atrophy, and depression in patients with MS [53]. The presence of depression was related to T1 lesions in the superior frontal and superior parietal regions. Depression severity was related to T1* lesions in the superior frontal, superior parietal, and temporal areas. Depression severity was also related to lateral and third ventricle enlargement and frontal atrophy. The authors postulated that depression in MS may be related to atrophy and disconnections between cortical and subcortical regions as a result of frontal and parietal destructive lesions in white matter.

INTERFERON-β THERAPY AND DEPRESSION

Initial suspicion of a link between IFN-β and depression in MS arose after reports of a suicide and attempted suicides during the first trial of IFN-β-1b [59]. Thyroid disease, including hyperthyroidism and hypothyroidism,
has also been linked to treatment with IFN-β [60]. Thyroid disease may masquerade as depression, and this possibility should be evaluated as part of the clinical assessment. Hence, for some patients, an association may exist between the use of IFN-β and symptoms of depression.

While the possibility of a link between depression and IFN-β exists and must be considered by the clinician, recent studies have not shown a clear association. Borras et al. reported findings from the first 2 years of a longitudinal study of MS patients treated with IFN-β-1b. At the onset of treatment, 90 patients were assessed, then 75 after 1 year and 56 after 2 years [61]. Worsening depression was not indicated in this study but rather a slight improvement in symptoms of depression and anxiety. Patten and Metz assessed depression in 365 patients with secondary progressive MS treated with either IFN-β-1a (Rebif®) or placebo [62]. No significant differences were noted between the two groups during 36 months of follow-up. Zephir et al. assessed depression in 106 MS patients treated with IFN-β-1a (Avonex®) [63]. At baseline, 85 percent of their patients had minimum or mild depression. Most of the patients (83%) retained their baseline status at the 12-month follow-up. Increased depression appeared to be correlated with increased disability. In contrast, the Controlled High-Risk Subjects Avonex® MS Prevention Study group, which evaluated IFN-β-1a therapy during a first demyelinating event, reported significantly more depression in the treatment group than the placebo group [64]. Hence, evidence to date from controlled studies does not establish a strong link between treatment with IFN-β and depression.

How to treat depression that occurs during IFN-β therapy is not settled. Once depression has been diagnosed, discontinuation of IFN-β is not the preferred option. Current evidence suggests that pharmacotherapy or psychotherapy for treatment of depression in MS, irrespective of its association to IFN-β, is a prudent strategy [65].

**IMMUNE DYSFUNCTION AND DEPRESSION**

Depression has been associated with an activated immune system. Compared with controls, depressed patients have increased numbers of activated T cells and acute phase proteins and higher production of proinflammatory cytokines, such as tumor necrosis factor (TNF-α), interleukin (IL)-6, IL-1β, and IFN-γ [66–70]. Lanquillon et al. found significantly decreased TNF-α levels in depressed patients who clinically responded to amitriptyline therapy compared with nonresponders [71]. This cytokine may be, therefore, a marker of psychopathological improvement.

MS is a chronic inflammatory demyelinating disease with predominantly T cell-mediated immune dysfunction. Like patients with depression, patients with MS have higher serum concentrations of proinflammatory cytokines [72]. Mohr et al. studied 14 patients with relapsing-remitting MS and major depressive disorder over 14 weeks [73]. They found that higher production of the proinflammatory cytokine IFN-γ by autoaggressive T cells in patients with MS was related to depression severity. Moreover, treatment of depression with both psychotherapy or pharmacotherapy may decrease IFN-γ production. Mohr et al. argue that the treatment of depression in patients with MS may not only provide mood stabilization but also a novel disease-modifying strategy.

Models that relate stress, cytokines, the endocrine system, and depression are supported in the literature [74–75]. In their literature review, O’Brien et al. noted that cytokines modulate corticotropin-releasing hormone, which produces heightened hypothalamic-pituitary-adrenal axis activity and increased adrenocorticotropic hormone and cortisol [74]. Whether these hormonal changes are primary or secondary pathological defects in the origin depression has yet to be determined. More research is needed for evaluation of the unique immune-endocrine relationships and their effect on mood in patients with MS.

**ASSESSMENT OF DEPRESSION**

Because depression is common and can temporarily fluctuate in patients with MS, healthcare providers must have the necessary tools to make timely and accurate diagnoses [62]. One difficulty in diagnosing depression in MS is that many depressive symptoms are observed in MS patients without mood disorders. For example, both fatigue and cognitive dysfunction associated with MS are common and must be considered by the provider in the differential diagnosis of depression. Although more rare, patients with MS can exhibit pathological laughter or crying that poorly correlates with the underlying mood. This syndrome of emotional lability typically occurs with bilateral forebrain disease [76]. Given the overlap of symptoms between depression and MS, the diagnosis of depression in patients with MS is often complicated. Therefore, researchers and clinicians have used psychometrically validated measures of mood to aid their differential diagnosis.
Historically, there have been two main theoretical frameworks for using validated instruments in the assessment of depression in people with MS. In the first framework, the main goal of assessment is to directly address the symptoms of depression. This is helpful when healthcare providers need to conduct rapid triage and diagnosis so that fast, directed treatment decisions can be made. The second framework looks at depressive symptoms in the context of other MS-related symptoms and emotional responses (e.g., fatigue, low self-esteem, anxiety) that affect overall QOL.

For the busy clinician, the U.S. Preventive Services Task Force recommends a brief two-question screening test for assessing depression (Figure 1) [77–78]. If the answer to one or both questions is positive, the patient should be assessed for the other seven DSM, Fourth Edition (DSM-IV) criteria for major depression. When using this two-question screening test, clinicians should recognize that some patients with MS demonstrate subsyndromal symptoms of depression or mood instability [79] that do not meet the criteria for major depression. These symptoms are nonetheless associated with significant distress and can respond to therapy. Thus, their detection with the two-question screening test would alert the clinician to the need for treatment. One should note that, despite its face validity, the two-question screening test is not yet validated for use with patients with MS. Thus, at present, it is preferable for clinicians to use an objective depression measure (discussed in the next paragraphs). Objective, validated self-report measures have been used in research on patients with MS, which provides at least some degree of validation. However, the two-question screening test may be useful for busy clinicians who have no time to administer objective measures and would otherwise forego more in-depth depression screening.

The BDI is one of the most commonly used objective self-report depression measures [50]. This inventory has a long history of proven validity for assessing depression in the general public. Though some researchers believe that use of BDI with patients with MS is valid [80], others believe that such use is inappropriate. One main argument against its use is the long administration time. For this reason, several studies have sought to develop questionnaires that assess depressive symptoms in a less time-intensive manner. An example is the Yale Single Question; however, it lacks sensitivity and failed to identify 35 percent of depressed MS patients who had been positively identified on the BDI [81]. Another argument against the use of the BDI with patients with MS is that, because of the high number of items that assess neurovegetative symptoms, it overdiagnoses depression in this population [15]. For that reason, a number of researchers have attempted to identify the BDI items that contain confounds between somatic MS symptoms and neurovegetative symptoms of depression. These studies have resulted in short-forms of the BDI [82]. The best validated BDI short form is the BDI-Fast Screen (BDI-FS). In a recent study, Benedict et al. validated the BDI-FS with patients with MS [83]. Data from the study supported the concurrent and discriminative validity of the BDI-FS, which indicates that the test did not confound MS-related neurological symptoms. The Chicago Multiscale Depression Inventory is an alternative short instrument that was developed for assessing depression in patients with MS [84]. It was found to have good internal consistency, sensitivity, and construct validity [4].

Based on the idea that a multifactorial approach is the most useful for obtaining a holistic picture of adaptation to MS, one movement in the MS field has been the development of a single multidimensional mood instrument. One such instrument is the Profile of Mood States (POMS),

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**Figure 1.**
which measures mood and multiple dimensions of adaptation, including daily activities, fatigue, and disease status [85]. Unfortunately, although the scale composite score has been found useful, the subscales appear to be intercorrelated [86]. Therefore, the effects of mood relative to other factors cannot be teased apart. A more recently developed multidimensional scale is the Functional Assessment of MS (FAMS). The reliability and validity of the FAMS in patients with MS have been confirmed [87]. The FAMS is divided into six subscales: emotional well-being (depression), mobility, symptoms, general contentment, thinking/fatigue, and family/social well-being. In their comparison of multiple QOL measures, Nicholl et al. found that the FAMS was the most sensitive measure of the functional domains often associated with decreased QOL in MS patients [88]. In a study that assessed QOL among patients with MS in Spain, MS disability was related to emotional well-being (depression), mobility, and physical symptoms [89]. This study confirms that although depression is certainly a factor in adjustment to MS-related disability, other issues must also be considered and assessed. At this point, little data exist on the FAMS with respect to more global use in the MS community.

Although the best multidimensional tool for assessing depression in patients with MS is currently unclear, the tool must effectively and independently measure both fatigue and depression. Data indicate that depressive symptoms can independently predict fatigue severity [90], and improvement in depressive symptoms appears to be closely associated with decreased fatigue severity [91]. This interaction between depression and fatigue has significant implications for QOL. Research on QOL indicates that the effects of depression on the overall well-being of patients with MS cannot be fully understood without consideration of the presence and severity of fatigue [92]. Given that both depression and fatigue may independently affect [93] and predict [94] QOL, how depression affects the well-being of patients with MS cannot possibly be understood without an evaluation of fatigue. One should note that, in addition to measurement of fatigue, a thorough multidimensional assessment of QOL in depressed MS patients should include, at minimum, a screen of cognitive assessment. In the last few years, the complex interactions among depression, fatigue, and cognition have been repeatedly described in the literature [95–97]. In fact, depressed MS patients clearly perform worse than nondepressed MS patients on measures of cognition [98]. This has significant implications for depressed MS patients with respect to their ability to function in their daily lives. When attempting to refine and expand multidimensional QOL measures, researchers must understand that depression does not exist in a vacuum for patients with MS. Specifically, for such instruments to effectively portray the effects of depression on an MS patient’s well-being, they must include items or scales that assess fatigue and cognitive function. Furthermore, multidimensional QOL scales should include some method for interpreting the interaction among depression, fatigue, and cognition. Only then can clinicians and researchers obtain a comprehensive picture of the QOL of depressed MS patients.

In sum, depression is a major problem facing patients with MS and has significant implications for QOL. For this reason, clinicians must perform a depression assessment. Figure 2 offers a model of depression assessment in a typical initial or follow-up clinical visit. For clinicians who do not have enough time to complete objective depression measures, the rapid two-question screening test (Figure 1) gives at least rudimentary information about depressive symptoms and the possibility of a DSM-IV diagnosis. Given that this screen has not been validated, however, a formal objective measure of depression that has been used in research on patients with MS is preferable. The most promising of these formal instruments is the BDI-FS; the Chicago Multiscale Depression Inventory also appears to be a useful unidimensional depression measure for patients with MS. Clinicians who complete a more comprehensive, multidimensional evaluation of depression will find that the results are more helpful in understanding their patients’ QOL. Following assessment, the clinician should determine whether further assessment (e.g., neuropsychological assessment) and/or treatment is required.

Multidimensional measures such as the POMS or FAMS offer useful information about mood and assess other domains that might interact with mood and affect patient well-being. Finally, recent research indicates that accurate assessment of the effect of depression on a patient with MS requires a scale that assesses fatigue and cognitive function. If this is not possible in the clinician’s office, referral to a neuropsychologist who specializes in MS may be warranted. A neuropsychological evaluation provides a thorough cognitive and psychiatric evaluation and an in-depth assessment of the interaction of depression, cognition, and fatigue.

SUICIDE AND MULTIPLE SCLEROSIS

Left untreated, MS and depression are unlikely to resolve and may potentially exacerbate each other. The
combination may ultimately lead to suicide. Investigators in Denmark showed that the cumulative lifetime risk for suicide from onset of MS was nearly twice that expected (standardized mortality ratio = 1.85) [99]. Suicide was the cause of 15 percent of deaths in a long-term follow-up study in Canada between 1972 and 1988 [100].

Feinstein evaluated risk factors for suicide in MS among patients of an outpatient clinic [10]. Severity of major depression, history of alcohol abuse, and living alone had an 85 percent predictive accuracy for suicidal intent. While all patients with MS should be assessed for depression and suicide risk, patients with these additional risk factors should be monitored closely. Male sex, onset of MS before the age of 30, and hopelessness are recognized as important predictors of suicide risk [101–102]. Individuals with suicidal ideation or self-injurious behavior require immediate psychiatric referral and possible inpatient admission.

**PSYCHOTHERAPY FOR DEPRESSION**

Psychotherapy has long been recognized as an important treatment for depression in MS. The earliest reports in the literature typically described group-based interventions that improved psychosocial functioning, such as depression, anxiety, understanding of physical limitations, and reduced use of medical services [103–104]. Information on inclusion criteria, treatment content, therapist adherence, treatment exposure, and outcome measures was often limited or absent.

**Group Cognitive-Behavioral Therapy**

The last three decades have seen a substantial increase in the quantity and rigor of empirical studies examining treatment of depression in MS. Variations on cognitive-behavioral therapy (CBT) are most frequently reported in the literature. Larcombe and Wilson presented the first results of a group-based CBT trial in patients with MS [105]. Patients were randomly assigned to six weekly sessions of traditional CBT or to a wait-list control group. CBT treatment focused on behavioral activation for increasing pleasant activity and social interaction, and on cognitive restructuring for identifying and challenging maladaptive thoughts and beliefs associated with depression. Large, statistically significant reductions in depression were noted across most outcome measures in the treatment group, while small increases in depression were observed in the control group.

In another study of group-based CBT, Crawford and McIvor examined the efficacy of a 13-session stress management program in patients with MS [106]. Individuals who received the intervention (n = 23) reported significantly lower levels of depression and anxiety at follow-up, whereas the control group (n = 21) reported increased symptoms.

Somewhat similar results were obtained by Tesar et al. who evaluated a seven-session psychotherapy group that emphasized relaxation training, development of cognitive and behavioral stress coping skills, and exercises
promoting body awareness [107]. Individuals were quasi-randomly distributed so that the first 14 consecutive patients visiting an MS outpatient clinic were assigned to the intervention group and the next 15 were assigned to the control group. Although depression in the intervention group was not significantly decreased relative to the control group, the use of depressive coping strategies was significantly improved.

**Individual Cognitive-Behavioral Therapy**

Using a psychological-symptom management program known as stress inoculation training (SIT), Foley et al. compared individuals randomly assigned to six sessions of individual SIT with a wait-list control group that received standard care that included at least some supportive counseling services [108]. Significant reductions in depression and anxiety and increases in problem-focused coping were reported in the SIT group but no corresponding changes were reported for the control group.

In recent years, psychotherapy researchers have increasingly emphasized the tailoring of interventions to the specific needs of patients with MS. Recognizing that physical impairments and fluctuating symptoms may affect a patient’s ability to attend clinic appointments and participate in therapy, Mohr et al. developed and evaluated a telephone-based CBT for depression. Individuals were randomly assigned to receive weekly, individual telephone sessions for 8 weeks or to receive usual care [109]. Outcome analyses demonstrated significantly reduced depression in the treatment group but no change in the control group.

**Other Psychotherapies**

A small number of studies have addressed the efficacy of other psychotherapy modalities. Crawford and McIvor evaluated an insight-oriented group psychotherapy for treating depression in MS patients [110]. Subjects were randomly assigned to either an insight-oriented treatment group that met twice a week for 25 weeks, a current-events group that met with the same frequency, or a no-treatment control group. Small but significant reductions in depression were noted in the treatment group as compared with the other two groups. Maguire described a six-session group therapy that involved a combination of relaxation training, visualization, and drawing [111]. Participants reported decreases in negative mood states from pre- to posttest, although comparisons between treatment and control groups were problematic.

In the most comprehensive, randomized, controlled trial of treatment for depression in MS, Mohr et al. examined the comparative efficacy of CBT, supportive-expressive group (SEG) therapy, and the antidepressant sertraline [112]. Individuals randomized to the CBT condition received 16 weeks of individual sessions that focused on developing general cognitive and behavioral coping skills and MS-specific management skills. Individuals randomized to the SEG group received 16 weeks of group-based psychotherapy. Significant reductions in depression were noted across all three treatment groups; however, CBT and sertraline more effectively reduced depression scores than SEG. The authors noted that this result opposes findings in nonmedical populations in which all treatments for depression are generally considered equal. This discrepancy supports the hypothesis that depression in patients with MS and possibly other autoimmune diseases is etiologically different from depression in the general population [113–115].

In summary, psychotherapy appears to effectively treat depression in MS. Both group- and individual-based treatments reduce depressive symptoms. CBT interventions that focus on specific coping skills and MS symptom management are generally more effective than interventions that emphasize emotional expression or knowledge. Moreover, CBT is as effective as antidepressant medication.

**MEDICATIONS**

Antidepressant medications effectively treat patients with depression and MS [116]. Schiffer and Wineman presented results of the first double-blind placebo-controlled treatment trial of major depressive disorder in patients with MS [117]. A total of 28 patients were enrolled; one half received a 5-week trial of desipramine and psychotherapy and the other half received a placebo and psychotherapy. As evaluated by the Hamilton Rating Scale for Depression and the BDI, patients treated with desipramine improved significantly more than patients in the placebo group. Anticholinergic side effects, however, were reported in a significant majority of desipramine-treated patients. In an open-labeled trial of 11 depressed patients with MS, Scott et al. demonstrated that sertraline was both effective and well tolerated [118]. Similarly, Mohr et al. demonstrated that a 16-week trial of sertraline significantly reduced depression scores in patients with MS and major depressive disorder [112].
A number of medical therapeutic options are open to MS patients with depression. Drug classes of major antidepressants and representative medications are listed in the Table. Tricyclic antidepressants are among the oldest and have a long track record of success in treating depression. Their anticholinergic and cardiac side effects are problematic for many patients as noted in the trial by Schiffer and Wineman [117]. In MS patients with significant neurogenic bladder symptoms, however, these anticholinergic effects may be helpful. Newer agents are well tolerated with some subtle differences in side effect profiles. In general, the effectiveness of antidepressants is comparable between and within classes of medications [119]. A particular agent should be chosen based largely on its side effect profile, safety, quality of clinical trial data, cost, and patient preference. The selective serotonin reuptake inhibitors (SSRIs) have been proven to be well tolerated, safe, and effective in patients with MS and are a good initial choice for pharmacotherapy.

A healthcare provider must conduct an adequate trial and assessment period to judge the success of an antidepressant. A stable dose of an individual drug should be given for at least 6 to 12 weeks. Before a medication is started, side effects must be reviewed with the patient and monitored regularly. Finally, response to treatment should be assessed with an easy-to-administer scale, such as the BDI or BDI-FS. A positive response to a particular antidepressant is typically defined as a 50 percent reduction in a standardized depression scale score. A nonresponse to medication is defined as no net change from baseline of the standardized depression scale score [120].

**OTHER THERAPIES**

In the general population, up to 70 percent of patients with major depression respond to an adequate antidepressant medication trial. For those who do not respond, alternative approaches are needed. Studies in the general population have shown that regular exercise effectively reduces depression [121]. In a controlled randomized trial, Petajan et al. showed that patients with MS had significantly reduced depression after 15 weeks of aerobic training [122]. For severe depression in MS, electroconvulsive therapy (ECT) has been shown to be useful [123–124]. Some concern remains, however, that ECT may mediate neurological deterioration by altering the number or size of central nervous system plaques [125]. Recent studies in non-MS populations have found that stimulants such as methylphenidate [126] and modafinil [127–128] are useful adjunctive treatments for depression. In general, when therapeutic medication failures are encountered, the dose, compliance, and duration of treatment should be reassessed. Alternatively, the healthcare provider should question the diagnosis and rule out potential depression masqueraders.

### Table.
Medication options for depression.

<table>
<thead>
<tr>
<th>Drug Class</th>
<th>Representative Medications</th>
<th>Dosing Range (mg/day)</th>
<th>Untoward Effects</th>
<th>Comments</th>
</tr>
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<tbody>
<tr>
<td>Norepinephrine Reuptake Inhibitor (tricyclic antidepressants)</td>
<td>Nortriptyline</td>
<td>50–200</td>
<td>Anticholinergic symptoms</td>
<td>Anticholinergic properties may help with neurogenic bladder symptoms</td>
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<tr>
<td></td>
<td>Desipramine</td>
<td>100–300</td>
<td>Lethargy</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Protriptyline</td>
<td>15–60</td>
<td>Cardiovascular effects</td>
<td></td>
</tr>
<tr>
<td>Selective Serotonin Reuptake Inhibitor</td>
<td>Sertraline</td>
<td>50–200</td>
<td>Insomnia</td>
<td>Well-tolerated first-line agents</td>
</tr>
<tr>
<td></td>
<td>Fluoxetine</td>
<td>20–80</td>
<td>Sexual dysfunction</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Citalopram</td>
<td>20–40</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>Selective Norepinephrine Reuptake Inhibitor</td>
<td>Mirtazapine</td>
<td>15–45</td>
<td>Somnolence</td>
<td>Minimal effect on sexual function</td>
</tr>
<tr>
<td></td>
<td>Nefazodone</td>
<td>300–600</td>
<td>Weight gain</td>
<td></td>
</tr>
<tr>
<td>Norepinephrine Dopamine Reuptake Inhibitor</td>
<td>Bupropion</td>
<td>150–450</td>
<td>Seizures, psychosis</td>
<td>Minimal effect on sexual function</td>
</tr>
<tr>
<td>Serotonin Norepinephrine Reuptake Inhibitor</td>
<td>Venlafaxine</td>
<td>75–225</td>
<td>Sustained hypertension, withdrawal syndrome</td>
<td>Dose titration needed</td>
</tr>
</tbody>
</table>
USE OF DEPRESSION GUIDELINES

Depression in patients with MS is commonly underdiagnosed. The reasons for this underdiagnosis likely parallel those found in general primary care settings where depression is also frequently underdiagnosed. A number of studies have shown that providers’ use of guidelines along with a thorough follow-up program in primary care settings can facilitate successful identification and treatment of patients with depression [129–130]. Two frequently cited guidelines for depression treatment were published in 1993 by the Agency for Health Care Policy and Research (AHCPR) [131] and the American Psychiatric Association [132]. A more recent update to the AHCPR guidelines focuses more on their use within primary care settings [133]. The authors found that antidepressant pharmacotherapy and psychotherapies are effective when transferred from the psychiatric to primary care environment. After discussing the options with the patient, the clinician should choose a therapeutic approach based on patient preference. Most importantly, studies have suggested that improved treatment of depression within primary care requires an organized treatment program, regular patient follow-up with monitoring of treatment compliance and, in more severe cases, a prominent role for the mental health specialist.

Despite studies that support guideline implementation, other reports have shown that guidelines fail to improve depression treatment [134–135]. Current evidence suggests that without a structured follow-up system, guidelines are unlikely to be implemented successfully. In many clinics, time and lack of resources often preclude the use of guidelines.

We would argue that the high frequency of depression in patients with MS makes periodic screening as outlined in Figure 1 a high-yield activity. Patients with MS should be screened at least annually for depression. Patients who meet the criteria for depression should be treated with a single antidepressant or psychotherapy based on the previously discussed results. Monitoring with the BDI, BDI-FS, or a similar assessment tool should be performed 4 to 6 weeks after initiation and periodically thereafter. Indications for referral to a mental health specialist should include severe symptoms, suicide risk, comorbid psychiatric or substance abuse disorder, and failure to respond to appropriate treatment.

Treatment algorithms have been developed that address alternative treatments and help the clinician make evidence-based choices. The Texas Medication Algorithm Project (TMAP) published a depression treatment algorithm that was recently implemented at a number of sites [136]. The TMAP has three major phases: acute (stage 1), continuation (stages 2–4), and maintenance treatment (stage 5). Patients move between stages based on inadequate symptom improvement or untoward effects from therapy. Another algorithm, called sequenced treatment alternatives to relieve depression (STAR*D), is under development [137]. STAR*D is based on a prospective study that randomly assigned patients who had not achieved remission with an SSRI to one of six treatment options. STAR*D and TMAP results regarding compliance, remission, and cost-effectiveness will be very useful for patients with depression and their healthcare providers. More research on the unique treatment issues of MS patients with depression and on the development of effective algorithms is needed.

CONCLUSIONS

Depression is the most frequent psychiatric disorder in MS patients. Its etiology is multifactorial and associated with focal demyelinating lesions and psychosocial stress. Proper diagnosis and severity assessment are critical prior to initiation of therapy. The method used for measuring depression in the MS population differs depending on the goals of the assessment. Patients with suicidal ideation should be referred for immediate psychiatric consultation and be closely monitored. While more therapeutic trials for depression in MS are needed, patients with MS have been shown to respond to current antidepressant medications. Unfortunately, many patients with MS and major depression or suicidal thoughts are under-recognized and undertreated. Unlike other aspects of MS, depression is treatable and is a potentially preventable cause of death. MS care providers need to vigilantly assess depression and suicide risk in their patients.

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