Relapses in Multiple Sclerosis: Definition, Pathophysiology, Features, Imitators, and Treatment

Multipl Skleroz Atakları Üzerine Güncelleme: Tanım, Patofizyoloji, Özellikler, Takliççiler ve Tedavi

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Summary

Relapse in multiple sclerosis (MS) is defined as a neurologic deficit associated with an acute inflammatory demyelinating event that lasts at least 24 hours in the absence of fever and infection. Myelinolysis and axonal transection occur in relapses. Diagnosis, prognosis, treatment, and many other features of the disease are directly related to the relapses. MS starts as the relapsing-remitting (RRMS) form in 85% of patients. A large number of relapses in the first years, polysymptomatic relapses, and pyramidal system, brain stem, and spinal cord involvement are signs of a poor outcome. The average frequency of relapses is approximately one per year during the first years of RRMS. The frequency of relapses increases during systemic infections, psychological stress, and in the first 3 months after birth. Seventy-five percent of relapses are monosymptomatic. Pseudo-relapses and paroxysmal symptoms are distinguished from relapses by their sudden onset, sudden termination, and shorter duration. Contrast enhancement is valuable in imaging, but undetectable in most relapses. The regression in the first few weeks of relapses is explained by reduction of the edema, and by remyelination in the following months. Relapses and their features are also among the main determinants of treatment. High-dose methylprednisolone and early treatment with adrenocorticotropic hormone reduce post-relapse disability and shorten the duration of relapses. Plasmapheresis is a good option for patients who do not respond to steroid treatment. Identification of relapses by patients and physicians, distinguishing them from imitators, proper evaluation, treatment when necessary, and monitoring the results are of great importance for patients with MS. The educational levels of patients and physicians regarding these parameters should be increased. Well-designed studies that evaluate the long-term effect of relapse treatment on disability are needed.

Keywords: Multiple sclerosis, relapse, definition, features, treatment

Öz


Anahtar Kelimeler: Multipl skleroz, atak, tanı, özellikleri, tedavi

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Received/Geliş Tarihi: 06.12.2015 Accepted/Kabul Tarihi: 10.04.2016

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Introduction

Multiple sclerosis (MS) starts as the relapsing-remitting (RRMS) form in 85% of patients (1). Therefore; one of the most significant features of clinical manifestations is usually the emergence of new symptoms or worsening of present symptoms within days, so called relapses. The diagnosis, prognosis, treatment, and many other features of MS are directly related to the relapses. In this article, the topics discussed will be briefly relapse pathophysiology, and mainly relapse definition, characteristics, and treatment.

Relapses are the main determinants of subtypes of MS. Approximately half of untreated patients with RRMS progress to SPMS at the end of 10 years (2). The determinants of progression to SPMS are the reduction in the frequency of relapses and progression between relapses. Although there is no consensus, there is another form of MS with a pattern of relapses within worsening neurologic functions from the onset of symptoms, namely progressive-relapsing MS (PRMS) (3,4).

The primary progressive form (PPMS) accounts for approximately 10-15% of cases. The suggestions of the last two years regarding subtypes of primary and SPMS forms were transformed into a definition and the active-inactive and progressive-stable subtypes were identified for both types. In this definition, the active and inactive forms are determined according to relapses and magnetic resonance imaging (MRI), and progressive or stable forms are based on disability score (5).

Prognosis in patients with MS is also directly related to relapses. The number and type of relapses within the first 5 years is important for the prediction of disease progression. Higher number of relapses in the first years, polysymptomatic relapses, and pyramidal system, brain stem and spinal cord involvement are poor prognosis indicators. Conversely, sensory relapses, fewer relapses, and relapses with optic neuritis and recovery without sequelae are good prognostic features (6,7).

Early treatment of relapses with anti-inflammatory drugs such as steroids or adrenocorticotropic hormone (ACTH) reduces post-relapse disability and shortens duration of relapses (8,9).

On the other hand; the efficacy of relapse treatment in preventing and reducing long term disability is not based on scientifically solid evidence. The lack of such studies can be explained by several reasons. First, such studies should be long-term studies so that they can achieve reliable results. In addition, besides relapse treatments, immunomodulators and immunosuppressants are used in almost all patients at different stages, with various medications and applications. Furthermore, relapse treatment is performed in all patients except for patients who refuse on their own requests and formation of a subgroup without relapse treatment is considered unethical in studies. One of the other main reasons is that the pharmaceutical industry promotes studies on treatments that aim to change disease course positively rather than studies on relapse treatments. The proper application of relapse treatment may be as important as immunomodulatory and immunosuppressive therapies.

The Pathophysiology of Relapse and Symptoms

MS relapses are characterized by following sequential events: Blood-brain barrier (BBB) disruption due to white matter lesions, immune reactive cell migration to this region, and damage to oligodendrocytes and myelin caused by these cells. Inflammation is primarily mediated by T-lymphocytes. The probable initiator of pathogenesis is the migration of autoreactive Th17 cells, first from peripheral blood into the cervical lymph nodes, then into cerebrospinal fluid (CSF), and then finally into the choroid plexus, meninges and parenchyma (10).

This is the first wave. After disruption of the BBB, more Th17 and Th1 cells invade the parenchyma in the second wave. Inflammation becomes evident with chemokines released by myelin-reactive T-cells. Changes in the expression of adhesion molecules on the surface of T cells and macrophages facilitate the transport of these cells across the BBB. Interleukin 1, 12, 17, and 23 have been demonstrated to play role in triggering inflammation (11). The second stage begins with migration of CD8+ T cells and more CD4+ T cells into the central nervous system (CNS). Inflammation activates microglial cells and activation of these cells is the main cause of degeneration in the early stage. Finally, myelinolysis, demyelination, and axonal transaction occur in MS relapse. Axonal transaction is associated with permanent disability. It is demonstrated that axonal transaction can begin in very early stages of a MS plaque, even in the 2nd week after its appearance in MRI (12).

The occurrence of symptoms in relapse depends on conduction blocks and delayed axonal conduction. Conduction velocity in the axolemma is associated with higher concentrations of sodium (Na) channels in the nodes of Ranvier. Na channel density significantly decreases in the axolemma located below areas of demyelination. Consequently, conduction blocks and significant reductions in conduction velocity occur, and classic MS symptoms emerge. According to another hypothesis, formed autoantibodies adhere to Na channels, which makes the axolemma non-excitable and causes conduction block (13). Demyelination areas should have a minimum diameter of 4 mm for the formation of conduction blocks in humans. These areas were determined to have a diameter of 2.5 mm in rats (14).

The release of nitric oxide by glial cells in lesions has been shown to directly slow axonal conduction. Inducible "nitric oxide
The Definition and Features of Relapse

The definition of relapse is mainly based on clinical findings. After the last revision, it was defined in the 2010 McDonald Criteria as “patient-reported symptoms or objectively observed signs typical of an acute inflammatory demyelinating event in the CNS, current or historical, with duration of at least 24 hours, in the absence of fever or infection” (16).

A neurologist should distinguish between pseudo-relapses and true relapses. Pseudo-relapses will be discussed under the title “Relapse imitators.” Relapse should be considered even if a previous symptom arises and lasts more than 24 hours, and especially if the emerging neurologic deficit is evident. Proper treatment according to relapse features should be considered because up to 70% of MS relapses are in the form of exacerbation of previous symptoms. In addition, almost all causes of pseudo-relapses, mainly infections, may also trigger true relapses. The triggering effect of menstruation alone on true relapses could not be demonstrated. However, there are findings showing worsening of symptoms in the pre-menstrual period (17).

It is of great importance that patients with MS inform physicians or health care personnel about possible relapse symptoms. In a recently published study, it was shown that relapses were under-reported to health care professionals, with 28% of respondents failing to report their most recent attack, and 46% declared that they had failed to report an attack in the past, even in England where MS prevalence and socio-cultural level are very high (18). In another study in patients with MS, the reporting time of relapses to MS specialist nurses and family physicians was found as 10 days and 58 days after onset, respectively (19). There have been no studies in this field in Turkey.

In previous treatment efficacy studies conducted between 1960-2008, the average frequency of relapses in placebo control groups was approximately one per year. The annual average frequency of relapses in the treatment groups was found as 0.68 in these studies (20).

In general, we know that the frequency of relapses is higher in women and at younger ages, and decreases with time. However, it is difficult to know whether this decrease is due to treatment of almost all of the patients on any occasion or to the natural course of the disease. Both may have an effect on the decrease in relapse frequency (21,22).

The three unquestionable conditions with increased relapse frequency are systemic infections, psychological stress expressed by patients, and the first three months after birth (21,22). In these conditions attack triggering is thought to be primarily due to changes in cytokines and nitric oxide. Other reasons are controversial. The frequency of relapses increases in the spring and slightly reduces during winter months. In the recent years, this seasonal relationship is postulated to be associated with serum vitamin D levels (23). Data coming from relevant studies show that lower vitamin D levels are associated with more frequent relapses (24,25). It has been demonstrated that previously accused physical trauma, vaccinations, and epidural anesthesia do not have relapse triggering effects. The reduction in relapse frequency was found partially related with higher serum levels of vitamin D and omega-6 fatty acids, and sunlight exposure (22).

A typical relapse reaches its peak within a few hours and days, and recovers partially or completely after a plateau period lasting days and weeks. Particularly in some of infection-triggered relapses, fluctuations may be observed or, increases or decreases in symptoms may interweave in polysymptomatic relapses. In this case, it is difficult to distinguish whether symptoms are related to a single relapse or multiple relapses (7). Therefore, there should be at least 30 days after any previous attack or a stable period to be considered as two separate relapses (26).

In addition, multiple relapses that emerge with multiple symptoms in a short period of time may mimic transformation to progressive phase, and vice versa. So, the progressive phase may be perceived as severe relapses with short intervals by patients and physicians. Although very rare, it should be noted that relapse might occur without an obvious physical finding. Relapse with fatigue and cognitive impairment may be observed (27,28,29). If there are multiple paroxysmal symptoms that last more than 24 hours, this is also described as relapse (30).

Although the phrase “there should not be a systemic infection” is included in the definition of relapse, we should not forget that about one third of MS relapses are triggered by systemic infections. The risk of relapse increases during the 2-week period before and until 5 weeks after the onset of severe infections (-2/+5 week rule) (21,31).

Most of relapses negatively affect activities of daily living, and increase emotional stress and depression. Quality of life is reduced in patients with frequent relapses. The psychosocial burden due to relapse has a negative effect on social life and the family life of patients (6).
The most comprehensive study ever made about MS relapse was published a short time ago. Fifty thousand relapses of approximately 15,000 patients were collected using the MSBase database and it was shown that three out of four relapses were monosymptomatic. The most common type of relapse was sensory relapse (48%). This was followed by pyramidal (34%), brainstem (22%), visual (20%), cerebellar (7%), sphincter (3%), and cognitive dysfunction (1.4%) relapses, respectively (32).

According to the data of another recent study that evaluated relapse symptoms from the patients’ perspective, fatigue was the most common (68%) relapse symptom expressed by patients. It was followed by leg-foot weakness (63%), numbness-tingling (55%), difficulty in walking (51%), hand-arm weakness (49%), coordination difficulties (47%), dizziness-balance (46%), muscle tightness-stiffness (46%), cognition (thinking-memory) problems (42%), vision problems (34%), bladder problems (31%), pain-burning-itching (27%), bowel problems (19%), speech changes (16%), chewing-swallowing problems (14%), and sexual problems (8.5%) (33). Visual, sensory, and brainstem relapses were found more frequently in early disease and in younger patients. Pyramidal, sphincter, and cerebellar relapses were more common in older patients and in progressive disease. Women presented more often with sensory or visual symptoms. Men were more prone to pyramidal, brainstem, and cerebellar relapses.

About 70% of new relapses present with clinical signs similar to previous relapses. However, the exact same presentation is rare. Different relapses with similar phenotypes are typical. This situation is explained by the presence of new demyelination regions on or close to former remyelination regions (32,34).

The data obtained in recent years show that patients with suspected relapse should be evaluated in terms of overlooked fatigue and cognitive impairment. More than half of patients (55.2%) stated that their school or work lives were impacted and it was shown that this duration was 12.7 days on average (33).

The average frequency of relapse in the early years of RRMS is approximately one per year. About 17% of untreated relapses recover with significant sequelae (35). Each relapse causes an average of 0.24 to 0.57 residual change in the Expanded Disability Status Scale (EDSS) score and full recovery after relapses were observed in only 55% of patients in terms of EDSS scores (36,37).

There is no scale used to determine the severity of relapse. However, a recently published study evaluated oral steroid administration in relapse. Relapses were identified as “severe” (increase >2.5 points on EDSS), “moderate” (increase 1-2.5 points on EDSS), and “mild” (increase ≤0.5 points on EDSS) (38).

Full recovery from a relapse is more difficult in polysymptomatic and spinal relapses, and in the elderly. Relapses presenting with pyramidal tract and sphincter disorders have a poor long-term prognosis when compared with relapses without involvement of these systems (39,40,41,42).

The prognosis in patients with insufficient regression after the initial relapse and patients who present with brainstem, cerebellum or spinal cord syndrome was worse. The mean time to progression to SSPM was 30.2 years in half of the patients with relapses recovered fully or close to fully within the first 5 years, whereas this period was 8.3 years in poor recoverers (43).

While the majority of improvement in disability occurs within the first 2-3 months in relapse, this period may sometimes even exceed 1 year. The regression in the first few weeks of relapses is explained by reduction of plaque and surrounding edema, and by remyelination in the following months (44).

- Relapse: Neurologic deficit associated with an acute inflammatory demyelinating event that lasts at least one day, in the absence of fever and infection.
- Up to 70% of MS relapses are in the form of exacerbation of previous symptoms.
- All causes of pseudo-relapses, except menstruation, may also trigger relapse.
- The average frequency of relapses detected in natural course studies is approximately one per year.
- The frequency of relapses is higher in women and at younger ages, and decreases with time.
- The frequency of relapses increases during systemic infections, psychological stress, and in the first 3 months after birth.
- A typical relapse reaches its peak within a few hours and days, and recovers partially or completely after a plateau period of days and weeks.
- Neurologic deficits need to occur at least 30 days after any previous episode or a stable period to be considered a new relapse.
- Seventy-five percent of relapses are monosymptomatic.
- The risk of relapses increases 2 weeks before and until 5 weeks after the onset of a severe infection.
- The most common relapse symptom expressed by patients is fatigue.
- Seventeen percent of untreated relapses recover with significant sequelae.
- Regression is more difficult in polysymptomatic, spinal, and advanced-age relapses.
- Regression in the first few weeks of relapse is explained by reduction of the edema, and by remyelination in the following months.

Relapse Imitators

Pseudo-relapse; is the emergence of previous symptoms accompanied by increased body temperature as a result of febrile diseases, menstruation, hormonal disorders, hunger, and fatigue. Symptoms occur as a result of reduction block in damaged axons due to temperature increase. There are no symptoms that were not present previously and they usually last less than a day. It is always a repetition of previous symptoms. Pseudo-relapse starts with the emergence of the cause and finishes with the disappearance. It usually disappears at rest and in cold environments.
Paroxysmal symptoms; tend to repeat and are temporary disorders such as tonic spasms, weakness, dysarthria, ataxia, trigeminal neuralgia, and Lhermitte and Uhthoff phenomena. If these symptoms last longer than 24 hours, they are qualified as relapse. Almost all of them start abruptly and end abruptly. The vast majority of them last seconds to minutes.

Daily changes can be monitored in patients, particularly in those with high disabilities, and they can be misinterpreted as relapse. The majority of these patients have no relapse expectations, and reductions and increases in symptoms are observed in an unexpected shorter time in MS relapse.

- Pseudo-relapse is the re-emergence of previous symptoms; they emerge in situations that cause increased body temperature, end abruptly with termination of these situations, and last less than 1 day.
- Paroxysmal symptoms start abruptly; end abruptly, usually last seconds, and there is no expectation of relapse.

**Imaging in Relapse**

Relapse is a clinical definition. Among all the features of MS, history and physical examination are foremost in the evaluation and management of patients. A neurologist can determine a relapse based on the patient's history and examination findings, even in the absence of MRI lesions. However, the importance of imaging in relapse is increasing in light of developments in the MRI field. The radiologic equivalent of a MS relapse is an acute demyelinating lesion. However, most MS lesions are silent. Only one out of 8-9 lesions causes clinical symptoms. Contrast enhancement in the plaque is important in the assessment of disease activity and demonstrates BBB disruption in the plaque region. Enhancement may be diffuse or ring-shaped. A portion of medium and large-sized plaques tend to show diffuse contrast enhancement initially, but then a changeover to ring-shaped enhancement in time. “Open ring” enhancement is typical for MS and is important in distinguishing plaques from tumors and abscesses (Figure 1).

The presence of contrast-enhancing lesions consistent with symptoms in imaging is valuable in terms of relapse. However, contrast enhancement cannot be detected in many MS relapses. Contrast enhancement often begins before symptoms and before the appearance of T2 lesions in routine imaging, and lasts an average of 4 weeks. In recent studies that used high-resolution MRI techniques, some lesions were reported to be detected in fluid-attenuated inversion recovery (FLAIR) sequences before contrast enhancement. In a recent study of five untreated patients with weekly-performed three-dimensional FLAIR MRI, it was shown that 11.2% of lesions were visible on FLAIR images before the appearance of contrast-enhancement and 12.5% enhanced before being apparent on FLAIR (45).

Subclinical disease activity is seen on MRI in many patients. However, contrast enhancement in patients with clinical relapse is more frequent than in patients with subclinical MRI activity (46,47,48,49).

In routine practice, the standard dose of gadolinium is 0.1 mmol/kg. Gadolinium is administered intravenously for at least 30 seconds. Contrast enhancement is evaluated on T1 sequences. T1 imaging should be performed in 5 minutes following contrast administration. One of the major problems in demonstrating contrast enhancement is using an insufficient amount of contrast medium in some MR centers. Contrast medium should be administered as a full dose and even a double dose if requested by a neurologist.

- Most MS plaques are silent.
- Contrast enhancement is valuable, but it is not detectable in the majority of relapses. It usually begins before symptoms and the appearance of T2 lesions, and lasts an average of 4 weeks.
- “Open ring” enhancement is typical for MS.

**Relapse Treatment**

The aims of relapse treatment are shortening the relapse period and preventing or reducing potential disabilities. Treatment of relapses with high doses of steroids and ACTH reduces post-relapse disability and shortens duration of relapses. Relapse treatment has secondary benefits if it is successful: It can help patients to change...
their perception that MS is an incurable disease, can strengthen the patient-physician relationship, and can improve the patient adherence to follow-up and treatment. The long-term effect of relapse treatment on disability is controversial. There are no multicenter studies with high numbers of patients creating Level I and II evidences. Although many new developments related to immunomodulatory and immunosuppressive treatments in MS have been witnessed in the last 25 years, relapse treatment has remained unchanged. Relapses are mostly treated worldwide with high dose steroids except for those presenting with pure mild to moderate sensory involvement and this treatment procedure is recommended in all guidelines.

- Treatment of relapses with intravenous methylprednisolone (IVMP) and ACTH reduces post-relapse disability and shortens the duration of relapses.
- High-dose steroids exhibit anti-inflammatory effect by both genomic and non-genomic mechanisms (dual effect).
- Maintaining oral therapy after IV administration does not provide additional benefit.
- Daily IV dosage is 500-2000 mg and the administration period is 3-10 days. It is usually administered in 100-250 mL of 0.9% NaCl or 5-10% dextrose solution for at least 60 minutes.
- The acute adverse effects of IVMP treatment are hyperglycemia, gastrointestinal intolerance, insomnia, euphoria, depression, metallic taste in the mouth, flushing, and infections.
- Blood pressure measurement, blood glucose, sedimentation, C-reactive protein, electrolytes analysis, and urinalysis should be performed prior to administration. Treatment should be delayed in the event of clinical signs or apparent laboratory findings of systemic infection.
- Treatment should be applied in hospital in pregnant women, children, and the elderly, and in the presence of hypertension, diabetes mellitus and peptic ulcers.
- Unresponsiveness to IVMP treatment can only be considered at least 10 days after the end of administration.
- Plasmapheresis is a good option for selected patients who are refractory to steroid treatment and who are predicted to have a permanent disability and IV immunoglobulin (Ig) treatment has no role in relapses.

Adrenocorticotropic Hormone and Steroids

After it was understood that MS was an inflammatory and autoimmune disease, anti-inflammatory drugs were introduced in MS. Steroids are still the most powerful anti-inflammatory drugs among all drug groups. Synthetic IV steroids, and rarely oral forms, are the most commonly used drugs in MS relapse. There is Level I evidence regarding accelerated clinical improvement in MS relapse provided by steroids (50,51,52).

Steroids exhibit an anti-inflammatory effect by both genomic and non-genomic mechanisms (dual effect). The classic genomic mechanism recognized after the discovery of synthetic steroids which gains functionality with the activation of steroid-specific cytosolic receptors. Adrenal gland stimulation by low-dose external steroid or ACTH administration is sufficient for the activation of these receptors. The second mechanism is the non-genomic mechanism, which was discovered later and functions only by administration of very high doses of steroids. Non-genomic mechanism disrupts ion transport by direct effects on the cell membrane and induces apoptosis in divided T-cells. The efficacy of the non-genomic mechanism has been shown in experimental autoimmune encephalomyelitis, which is the experimental model of MS. Genomic and non-genomic mechanisms together compose the positive effects of steroids in MS relapse (53).

Another effect of corticosteroids is prevention of migration of inflammatory cells into the CNS. They also reduce synthesis of intrathecal immunoglobulin G (IgG), myelin basic protein, adhesion molecules, matrix metalloproteinases, and neurodegeneration products (54,55,56). All these biochemical findings are indicators of positive cellular and molecular effects of steroids on relapses. Steroid administration also reduces contrast enhancement. This is believed to occur due to the repairing effect of steroids on BBB. The contrast enhancement reducing effect of steroids lasts up to 7-9 weeks (57).

The first drug used in relapse is ACTH, which has an endogenous steroid-inducing effect (58).

They are rarely used nowadays because they do not increase plasma levels of steroids as much as high-dose MP, they act later, and are more expensive. Additionally, daily administration of 500-1000 mg of MP increases serum steroid peak levels 5-10 times more compared with ACTH administration (54). Despite these, positive effects equivalent to MP have been reported with ACTH use in relapse treatment (59,60,61). This effect is attributed to the direct anti-inflammatory and immunomodulatory effects of ACTH through melanocortin receptors. ACTH is showed to be effective in opsinclonus-myoclonus and infantile spasms via the melanocortin receptors whereas synthetic steroids are found to be ineffective in these conditions (62,63,64).

In practice, ACTH is administered 80 units/day intramuscularly or subcutaneously for 5-15 days. It has similar adverse effects to synthetic corticosteroids, but some patients who cannot tolerate MP treatment have been shown to tolerate ACTH treatment. It can be evaluated in patients who cannot tolerate high-dose MP therapy.

The most widely used drug in relapse treatment is MP. MP is a synthetic derivative of cortisol. It has a longer duration of action compared with cortisol, and it has a greater glucocorticoid effect but lesser mineralocorticoid effect. The half-life of circulating MP is 1.5 hours and the half-life of its metabolites is 4 hours. It reaches its peak concentration in 2 hours in plasma and 6 hours in CSF (65).

Low-dose oral maintenance therapy after IV administration is a preferred method sometimes. However, its superiority over
IV administration alone could not be demonstrated (66). In an international optic neuritis study, administration of 1 g/day IV MP for 3 days followed by 21 days of oral MP has been shown to be superior to oral MP administration alone. In another randomized, controlled, prospective study, EDSS scores were shown to significantly improve in steroid-treated groups in the 3rd and 6th weeks (67). Two different meta-analyses also reached similar conclusions. In one of these, 377 participants from six trials were evaluated and EDSS scores in patients treated with MP or ACTH were found significantly better at the end of 1 month (relative odds ratio: 0.37) (68). In another meta-analysis, similar results were achieved in terms of disability scores (69).

However, in another meta-analysis of studies that comprehensively assessed treatment of optic neuritis relapses with steroids, long-term beneficial effects of high-dose steroid use on disability could not be shown (70).

Administration of high doses of oral MP during relapse has been much debated in recent years. MP was used intravenously in the majority of the studies, but was also administered orally in some. No difference was found between groups regarding drug efficiency with administration of equal doses of oral and IV MP. In a well-designed study in which oral (1250 mg/per day for 3 days) and IV (1000 mg/ per day for 3 days), MP therapies were compared in 49 patients; no differences were found between the groups in the 1st, 4th, and 12th weeks in terms of both reducing EDSS scores and MRI lesions (71). In some centers, oral steroid administration is increased in patients, particularly in those who do not require close monitoring. However, oral MP is available only in the 16 mg form in Turkey and the patient needs to take 60-70 tablets every day. If oral administration is preferred, 25% higher dose than IV administration may be appropriate due to loss in the gastrointestinal tract.

The Adverse Effects of Steroids

In animal studies, oligodendrocyte-mediated remyelination has been shown to decrease with steroid use (72). Results supporting increased brain atrophy with steroids are controversial. This is more likely the “pseudo-atrophy” due to anti-edema effect. Steroids are also known to exhibit adverse effects on cognition and memory. However, these effects are almost always temporary. Bone formation immediately decreases and bone resorption increases with high-dose steroid. However, this effect is also probably temporary. A negative long-term effect of high-dose steroid administration during relapses could not be demonstrated (73).

Hyperglycemia and glycosuria (5%), gastrointestinal intolerance and dyspepsia, psychiatric effects such as insomnia, depression, and euphoria, and more rarely a metallic taste in the mouth, facial flushing, weight gain, paresthesia, and infections are among the other adverse effects of steroids. It can very rarely cause aseptic femoral head necrosis and cataracts. Acyclovir or valaciclovir can be used together if there is previous varicella-zoster infection in patients taking immunosuppressant drugs (74).

It is possible to come across different information regarding steroid administration in relapses when viewed from the patient’s perspective. In a study addressing this issue, one third of patients reported no adverse effects related to treatment and two-thirds reported one or more adverse events. Although one quarter of patients who reported adverse effects stated that these effects were moderate to severe, only 1% reported that they had to quit treatment because of the adverse effects. The most frequently reported adverse effect is insomnia (72%), followed by depressed mood (62%), metallic taste in the mouth (59%), headache (59%), anxiety (56%), and swelling of the body (52%) (20).

The Intravenous Administration of Methylprednisolone in Practice

Daily IV dose is usually 1 g and the administration period is 3-10 days. It is usually administered in 100-250 mL of 0.9% NaCl or 5-10% dextrose solution for at least 60 minutes. NaCl solutions and dextrose solutions are preferred in patients with diabetes and hypertension, respectively. There are also physicians who prefer 500 mg and 2 g as daily doses. In a study that compared the daily administration of 2 g and 500 mg IVMP, 2 g MP was found to be more effective in reducing the number of MRI contrast-enhanced lesions, but there were no differences regarding other measures (75).

Steroids are usually given in the morning so as to conform to the biorhythm. However, in a small study that compared IV steroid administration during the day or night in 17 patients, nighttime administration was reported to be better tolerated (76).

Blood pressure measurement, fasting blood glucose, sedimentation, C-reactive protein, electrolytes analysis, urinalysis, and chest X-ray should be performed prior to IV administration of MP. Treatment should be delayed in the event of clinical signs or laboratory findings of a systemic infection. If there are signs of asymptomatic urinary tract infection in laboratory findings, administration could be performed in conjunction with antibiotics.

If there are no contraindications, all patients should be given H2 receptor antagonists before administration and patients should be monitored for epigastric burning and dyspepsia. If clinical findings related to increased blood pressure emerge, patients should also be monitored at frequent intervals in this respect. The administration period should be longer in patients with cardiac diseases. Salt-free diet should be followed during administration.

Hypokalemia is the most common electrolyte disorder during administration. It should be considered in the presence of heart rhythm abnormalities and sudden weakness. The treatment should be applied in the hospital in pregnant women, children, and the elderly, and with hypertension, diabetes mellitus, and peptic ulcers.

IV administration can be applied in the 2nd and 3rd trimester of pregnancy, but should not be performed in the first trimester. Administration should be performed after milking the postpartum lactating mother and milk should be kept in the refrigerator. Mothers can breastfeed 4 hours after the end of administration. The baby can be fed with pre-milked milk during these 4 hours.

It is important to start high-dose steroids in the early stages (within one week) of relapse. If there is no contraindication, administration can be started on the same day in patients with severe relapse detected in the afternoon, and administration can be moved to morning hours with each new day.

The effect of high-dose steroid administration on long-term disability during relapses is controversial. There are no prospective studies in this regard and it is difficult for retrospective meta-analyses to truly reflect long-term disability. On the other hand, the
superiority of high-dose steroid administration over placebo was shown in terms of disability and the evidence was Level I. High-dose steroid administration during relapses causing disability is recommended in all guidelines.

Plasmapheresis

Plasmapheresis and IVIg trials have begun with a better understanding of the role of humoral immunity in MS pathogenesis. Data regarding the efficacy of IVIg in relapse is not clear. Despite positive results in early studies, the results from later studies were negative. Although it is specified as a 2nd choice in the European Federation of Neurological Societies guidelines, it has not included in America’s guidelines. In a study in 76 patients treated with either IVIg or placebo in addition to steroids, there were no differences between the groups (77). Today, it has no place in relapse treatment. The situation is different for plasmapheresis and it has up to 70% efficiency in relapses refractory to steroids.

In a group of 36 patients, mostly patients with MS, with relapse who had been diagnosed as having demyelinating disease and who failed to recover after treatment with IV corticosteroids, 42% and 6% improvement in neurologic disability occurred in the group treated with plasmapheresis and in the placebo group, respectively (78). In three retrospective studies in a total of 87 patients with definitive diagnosis of MS and refractory relapses to high-dose steroids, it was reported that effective treatment was achieved with therapeutic plasma exchange in two-thirds of patients (79,80,81). The situation is different for acute isolated optic neuritis refractory to high-dose steroid and no benefit from therapeutic plasma exchange could be demonstrated. However, it was found to be effective in optic neuritis accompanying neuromyelitis optica (82).

In patients with clinically isolated syndrome, acute relapse or acute worsening of symptoms, first-line treatment with therapeutic plasma exchange was found to have a 72% efficacy rate in a retrospective analysis. A median of 0.75 points reduction was achieved in EDSS scores (83). Another study showed that plasma exchange therapy was effective in relapse in SPMS; however, it had a low level of evidence (84). Unresponsiveness to steroid treatment in relapse can only be considered at least 10 days after the end of administration.

Plasmapheresis is performed via a central or peripheral venous catheter. A 1-1.5 x plasma volume is removed at each procedure every other day. A total of 5-7 exchanges are performed. It should not be performed in the event of active infection, hypotension, bradycardia, dehydration or leukopenia. The risk of serious complications is 4.6% and mortality risk is about 1.5%. Serious complications include anaphylaxis, thrombosis, sepsis, pneumothorax, hypotension, hyperthermia, hypokalemia and hypocalcemia. Good oral fluid intake should be encouraged on exchange days and the day before exchange to prevent hypotension.

Good prognostic features before plasmapheresis are short duration of illness, preserved deep tendon reflexes, and presence of surrounding edema and ring-shaped contrast enhancement on MRI (79). As a result, plasmapheresis should be performed in selected patients because of serious complications. It is a good option for relapses that are refractory to steroid treatment and that are predicted to cause permanent disability.

Although there is not enough scientific evidence available, high-dose cyclophosphamide or rituximab can be evaluated in patients who are refractory to steroid and plasma exchange therapy or in those for whom these therapies cannot be performed (85,86).

Conclusion

MS relapses and their features are among the main determinants of clinical picture and treatment. Identification of relapse by patients and physicians, distinguishing them from imitators, proper evaluation, treatment when necessary, and monitoring the results are of great importance. The educational levels of patients and physicians regarding these parameters should be increased. Well-designed studies that evaluate the long-term effect of relapse treatment on disability are needed.

Ethics

Peer-review: Externally peer-reviewed.

References


