The state of multiple sclerosis: current insight into the patient/health care provider relationship, treatment challenges, and satisfaction

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Background: Managing multiple sclerosis (MS) treatment presents challenges for both patients and health care professionals. Effective communication between patients with MS and their neurologist is important for improving clinical outcomes and quality of life.

Methods: A closed-ended online market research survey was used to assess the current state of MS care from the perspective of both patients with MS (≥18 years of age) and neurologists who treat MS from Europe and the US and to gain insight into perceptions of treatment expectations/goals, treatment decisions, treatment challenges, communication, and satisfaction with care, based on current clinical practice.

Results: A total of 900 neurologists and 982 patients completed the survey, of whom 46% self-identified as having remitting-relapsing MS, 29% secondary progressive MS, and 11% primary progressive MS. Overall, patients felt satisfied with their disease-modifying therapy (DMT); satisfaction related to comfort in speaking with their neurologist and participation in their DMT decision-making process. Patients who self-identified as having relapsing-remitting MS were more likely to be very satisfied with their treatment. Top challenges identified by patients in managing their DMT were cost, side effects/tolerability of treatment, and uncertainty if treatment was working. Half of the patients reported skipping doses, but only 68% told their health care provider that they did so.

Conclusion: Several important differences in perception were identified between patients and neurologists concerning treatment selection, satisfaction, expectations, goals, and comfort discussing symptoms, as well as treatment challenges and skipped doses. The study results emphasize that patient/neurologist communication and patient input into the treatment decision-making process likely influence patient satisfaction with treatment.

Keywords: health care provider survey, multiple sclerosis, patient-health care provider relationship, patient survey, treatment expectations, treatment satisfaction

Introduction

Multiple sclerosis (MS) is usually diagnosed in the third and fourth decades of life and can lead to fixed and/or progressive neurological impairment with a significant impact on activity.¹,² The last two decades have seen the advent of both disease-modifying therapies (DMTs) and new symptomatic therapies, adding to the complexity of the decision-making process of DMT selection and MS symptom management.³,⁴ More emphasis is being placed on patient-reported outcomes to help measure clinical outcomes, capture the full impact of the disease, and to determine health care provider (HCP) reimbursement.⁵,⁶ Thus, effective communication between patients and their
HCPs becomes even more important for understanding the full impact of the disease to achieve optimal clinical outcomes.

Shared treatment decision making is promoted as representing good practice, and likely encourages treatment adherence and improves patient satisfaction. To be active participants in the DMT decision-making process, patients need to have adequate understanding of both their disease and the benefits/risks of treatments. Patients with MS who are well informed about their disease have significantly higher medication adherence than those who are not. The key element to maintaining treatment adherence in patients with MS is an open and honest relationship between the patient and HCP. Determining perceived differences in communication effectiveness between patients and their neurologists is essential for improving clinical outcomes and quality of life.

The aim of this study was to assess the current state of MS care from the perspective of both patients with MS and neurologists treating patients with MS, and to gain insight into perceptions of treatment expectations/goals, treatment decisions, treatment challenges, communication, and satisfaction with care, based on current clinical practice.

Methods
Survey design
Closed-ended questions for an online survey were developed to assess patient and neurologist attitudes, perceptions, and knowledge related to different aspects of MS, including the impact of disease on patients’ lives, patient and HCP relationships, availability and usefulness of MS informational resources, and the expectations for satisfaction and also challenges encountered with a chosen DMT for MS. The survey was designed from a market research perspective for hypothesis-generating purposes. Questions were developed by the State of MS Consortium, comprising representatives from Biogen, GCI Health, and a steering committee (authors of this manuscript) of international MS experts and patient advocacy representatives, in collaboration with Harris Poll. Parallel questions for patients and neurologists were designed to allow comparisons. The survey responses were single or multiple response, numeric text, or rated on a 4-to-5-point Likert scale (eg, “very important”, “important”, “somewhat important”, or “not at all important”; “strongly agree”, “somewhat agree”, “somewhat disagree”, or “strongly disagree”, respectively). As the nature of this online survey was blinded and non-interventional, formal review and approval by the ethics review committee of the Hospital Vall d’Hebron was not required for this study. The study was carried out in accordance with the principles outlined in the Declaration of Helsinki.

Selection and recruitment
Neurologists and patients from Italy, Germany, Spain, the UK, and the US were surveyed between March 18 and April 25, 2014.

Patients
Recruitment targets were set at 100–500 patients per country and recruitment continued until targets were met. Patients (≥ 18 years of age) who participated in the research were recruited from survey panels provided by Harris Poll and vetted partners. Those invited to participate from the panels had previously agreed to participate in survey research, completed a double opt-in process, and identified themselves as having MS. Participating panel members were originally recruited into the survey panels using multiple methods (eg, directly, via banner ads, through other research). For this study, panelists were invited to participate in the research via email invitations. As access to the study was closed in each country once the recruitment target was met, determining the responder rate was not possible. A total of 2,023 panelists began the survey, with the 982 who completed it forming the analyzed cohort. Of those 1,047 invited panelists who started the survey but did not complete it, only 78 were determined to have met the age and diagnosis qualification criteria. The survey closed once the recruitment targets were met, so the responder rate could not be determined. Completion of the online survey questions deemed the qualify patients consented to participate in this study.

Neurologists
For neurologists treating patients with MS, recruitment targets were set at 150–200 neurologists per country and recruitment continued until the targets were met. US neurologists were randomly selected from the database of the American Medical Association master file. Those selected received a postal mail letter inviting them to participate in the research and were provided a link and password to the online survey. European neurologists were originally recruited to survey panels in a manner similar to patients from survey panels provided by Harris Poll and vetted partners and invited to participate in this survey via email. Neurologists had to have seen ≥ 1 patient with MS, on average, each year in their practice to be included in the survey. A total of 1,124 participants began the survey, with 900 who completed it forming the analyzed cohort. Of those 224 invited participants who started the survey but did not complete it, 80 were determined to have met the inclusion criteria. The survey closed once the recruitment targets were met, so the responder rate could not be determined. Completion of the online survey questions deemed the qualify neurologists consented to participate in this study.
For both neurologists and patients, specific subcategory quotas (e.g., MS type, years in practice) within each country and audience were not predefined; thus, baseline demographics may have varied by country.

Statistical analysis
In the US, patient results were weighted for sex, age, region, race/ethnicity, education, and income to align with the US MS patient population based on predetermined targets. The targets were based on a demographic profile derived from the polling company. Neurologist results were weighted for sex, region, and years in practice. US neurologist results were weighted to be representative of the American Medical Association neurologist population. No other countries were weighted individually because information on the population profiles was not available. Additional weights were applied to the total data to give each country equal representation in the overall results.

Descriptive statistics are reported for the total patient/neurologist samples. Given the aim of the project was to identify themes that might justify further study, no correction was made for multiple comparisons. Statistically significant differences across the subgroups of interest for patients or neurologists were determined by an independent sample t-test of column proportions and means at the 95% confidence level. No statistical testing was performed across audiences because the question phrasing differed between patients and neurologists. The dataset results were reviewed and any trends and patterns for themes of interest have been presented.

Results
Demographics
Nine hundred neurologists (Table S1) and 982 patients (Table 1) completed the survey.

| Table 1 Patient demographics |
|-----------------------------|
| Characteristics             | Germany | Italy | Spain | UK    | US    | Total |
| Patients (n)                | 146     | 100   | 100   | 125   | 511   | 982   |
| Type of MS (self-reported), % |         |       |       |       |       |       |
| RRMS                        | 43      | 40    | 42    | 41    | 66b-c | 46    |
| SPMS                        | 26a     | 35a   | 42a-c | 26c   | 18    | 29    |
| PPMS                        | 11      | 10    | 9     | 17a   | 8     | 11    |
| PRMS                        | 13c     | 5     | 4     | 6     | 4     | 6     |
| Other/not sure              | 7       | 10    | 3     | 10    | 5     | 7     |
| Mean (SD) age (years)       | 44 (10)c | 42 (12)| 40 (10)| 49 (13)d-e | 50 (13)d-e | 45 (12) |
| Female, %                   | 71a     | 59    | 44    | 74a   | 69a   | 72a   |
| Mean (median) time since MS diagnosis (years) | 11.8 (10)b-c | 8.6 (5) | 6.9 (5) | 12.3 (10)b-c | 14.3 (12)b-c | 10.8 (8) |

Notes: Weights were applied to the total data to give each country equal representation in the overall results. aSignificantly different than Germany. bSignificantly different than Italy. cSignificantly different than Spain. dSignificantly different than the UK. eSignificantly different than the US.

Abbreviations: MS, multiple sclerosis; PPMS, primary progressive multiple sclerosis; PRMS, progressive relapsing multiple sclerosis; RRMS, relapsing-remitting multiple sclerosis; SD, standard deviation; SPMS, secondary progressive multiple sclerosis.

Patients
Patients surveyed most commonly identified themselves as having relapsing-remitting MS (RRMS; 46%), followed by secondary progressive MS (SPMS; 29%) and primary progressive MS (11%).

Baseline characteristics differed by country. More participating patients in the US self-identified as having RRMS than other countries, while more patients in Europe self-identified as having any form of progressive disease compared with the US (Table 1). Patients from the US and UK were older, on average, than patients from Germany, Italy, and Spain and subsequently reported a longer average time since MS diagnosis (Table 1). Overall, 62% of patients had at some stage taken a DMT for MS (63% for RRMS, 68% for SPMS, and 52% for all others, including primary progressive MS, progressive relapsing MS [PRMS], and other/those who were not sure).

Relationship and communication with HCPs
The majority of patients somewhat/strongly agreed that their neurologists were honest in setting expectations for therapy success (85%), had a good understanding of all aspects of MS (84%), and had an open dialogue with them (82%). Although most patients (83%) reported that they were somewhat/very comfortable speaking with the neurologist or most frequently seen HCP about MS, nearly 1:5 patients reported being uncomfortable speaking about difficulty walking (19%), tremors (19%), and muscle spasms (18%; Table S2). Sensitive symptoms such as sexual difficulties (54%), bladder/bowel problems (28%), mood swings (26%), difficulty swallowing (21%), and cognitive/memory issues/trouble concentrating (21%) emerged as the most
uncomfortable for patients to discuss. Only 2%–3% of neurologists identified symptoms such as difficulty in walking, tremors, and spasms as uncomfortable topics for their patients; however, they may have overestimated patients’ discomfort talking about those same sensitive symptoms, including sexual difficulties (87%), bladder/bowel problems (54%), mood swings (37%), and cognitive/memory issues/trouble concentrating (37%; Table S2).

Communication appeared to be important in treatment satisfaction, as those who were satisfied with their DMT were more likely than those who were less satisfied to say that they were very comfortable speaking to their neurologist (64% vs 43%, respectively; \(P<0.05\)) and were more likely to say they had an open dialogue with their HCP in which they could ask any questions they wanted (92% vs 80%, respectively; \(P<0.05\)). The majority of patients somewhat/strongly agreed that their neurologist was accessible when needed (71%).

Cognitive impairment
More neurologists (74%) than patients (44%) felt MS has had a negative impact on the patients’ ability to perform cognitive tasks (Figure S1). When asked about MS symptoms they had experienced, 39% of patients reported cognitive/memory issues/trouble concentrating (Table S3). Thirty-seven percent of neurologists perceived their patients were not comfortable discussing cognitive impairment, whereas only 21% of patients were not comfortable discussing the problem with their neurologist (Table S2). While nearly all neurologists (95%) felt maintaining cognitive function was a very important/important part of their patients’ treatment goal, only 41% of patients who were currently taking DMTs reported it as part of their treatment goal.

Treatment selection, satisfaction, and expectations
Thirty-seven percent of patients reported that their HCP was the sole/primary DMT decision maker (Figure 1A), despite 71% of neurologists indicating that most patients with MS prefer that they, the neurologist, decide which treatment is appropriate for them. The level of patient participation in the treatment decision-making process remained constant across MS disease subtypes (Figure 1A). Patients in Spain (n=55) were the most likely (42%) to say their HCP was the sole decision maker compared with only 9% in the UK (n=47), 10% in the US (n=353), 14% in Italy (n=63), and 15% in Germany (n=125; Figure S2). Country-level results with small base size should be interpreted as directional in nature. Patients who were satisfied with their DMT were more likely to say that their expectations were very closely met than those who were not satisfied (67% vs 44%, respectively; \(P<0.05\)). More than half of patients who were currently taking DMTs (58%) were very satisfied/satisfied and 42% were somewhat/not at all satisfied (Figure 1B), but this varied by country. US patients were the most likely to be satisfied with their MS medication(s) and patients in Italy less likely to be satisfied than patients in most other countries (Figure S3). Most patients (69%) were optimistic about their future, and these patients were more likely to say they were very satisfied/satisfied with their current DMT (66%) than those who were not optimistic (37%). Patients who self-identified as having RRMS were more likely to be very satisfied with their current DMT than those who identified as having SPMS or other types of MS (31% vs 17% and 16%, respectively; \(P<0.05\)). Patients who were satisfied with their DMT were more likely than others to report having been an equal partner or primary/sole decision maker in choosing their DMT (69% vs 58%, respectively; \(P<0.05\)). Most neurologists (92%) agreed that MS patients who provided input when choosing a treatment plan tended to be more satisfied with their treatment experience than those who did not.

Only approximately half of patients (53%) said that their DMT met the expectations set forth before treatment closely/very closely. More than 1:5 patients either said no expectations were set (12%) or they could not recall if they were set (10%).

Treatment choice and goals
Both patients and neurologists included safety and side effects as top factors that are important when making decisions about DMTs (Figure S4). Patients’ top treatment goals for their current DMT included reducing relapse frequency (61%), slowing disability progression (60%), feeling better (53%), and reducing relapse intensity (50%). Similarly, neurologists’ most important treatment goals for their patients’ current MS DMT were reducing relapse frequency (97% very important/important), slowing disability progression (97%), feeling better (92%), and reducing relapse intensity (93%).

Treatment challenges
Top challenges identified by patients in managing their DMT were cost, side effects/tolerability of treatment, and uncertainty if treatment was working (Figure 2). Personal cost was identified as a challenge by more patients in the US than in Germany and the UK (36% vs 23% and 9%, respectively). Patients and neurologists were aligned in identifying top challenges, with the exception of cost, but differed in the frequency of their response (Figure 2). However, approximately half of neurologists also cited lack of acceptance of the disease, while only a few patients themselves reported this.
Nearly half of patients (49%) reported never skipping doses of their DMT. Of the 51% of patients who reported skipping doses, only 68% told their HCP when they skipped a dose. The top reasons patients cited for skipping a DMT dose were forgot (43%), side effects (22%), and feeling better (15%; Table 2). However, neurologists believed that side effects (73%) were the main reason for skipping doses.

**Skipped doses**

Nearly half of patients (49%) reported never skipping doses of their DMT. Of the 51% of patients who reported skipping doses, only 68% told their HCP when they skipped a dose.

**Figure 1** (A) Participation in DMT decision making and (B) satisfaction of patients with DMT.

**Notes:** (A) Qualified patients with MS who had taken prescription medications responded to the question, “How much input did you have in choosing your current/past disease-modifying medication(s), compared to your health care provider?” (B) n=463 patients with MS who were currently taking prescription medication responded to the question, “Overall, how satisfied are you with your current disease-modifying multiple sclerosis medication(s)?” The other category includes PPMS, progressive relapsing multiple sclerosis, and other/not sure. *Indicates RRMS significantly different from SPMS and other, or SPMS and other significantly different from RRMS (P<0.05). *Please note small base size; results should be interpreted as directional in nature. Total may exceed 100% due to rounding.

**Abbreviations:** DMT, disease-modifying therapy; HCP, health care provider; MS, multiple sclerosis; PPMS, primary progressive multiple sclerosis; RRMS, relapsing-remitting multiple sclerosis; SPMS, secondary progressive multiple sclerosis.
Discussion

This large global online survey evaluated MS patient and neurologist attitudes about treatment expectations and goals, treatment decisions, treatment challenges, communication, and satisfaction with care. Overall, patients and neurologists identified similar treatment goals and challenges. Patients who were satisfied with their current DMT were especially optimistic about the future. Patient input into their DMT decision-making process influences satisfaction, as does open communication, according to both patients and neurologists.

These results are consistent with another study that showed 79% of patients with MS preferred to take an active role in medical decision making. A shared medical decision-making process regarding DMT choice is important in MS because current therapies are disease modifying, not curative, and adherence is important to optimize clinical outcomes. Studies have suggested that shortcomings in communication and information provided by HCPs may affect the quality of care, which in turn may affect treatment adherence, clinical outcomes, and ultimately quality of life. While high levels of satisfaction were noted in many aspects of MS management, important gaps were identified. Key differences exist in the perceptions of patients and neurologists when discussing MS symptoms. Neurologists tend to overestimate their patients’ discomfort talking about difficult topics such as sexual difficulties, bladder/bowel problems, and cognitive impairment, but tend to underestimate their discomfort discussing topics such as tremor and walking difficulty. These discrepancies could potentially lead to neurologists failing to identify and address modifiable symptoms. Cognition has been shown to be key in patients’ quality of life and therefore it is a very relevant symptom. Cognitive impairment was of lower importance to patients than relapses and disability progression in most responses. The low number of patients identifying cognition as a problem may be related, at least in part, to the lack of correlation between perceived and objective cognitive issues, a disparity which has been previously noted. However, because cognition was not assessed in this survey, we cannot rule out that a higher proportion of patients without cognitive problems completed the survey.

Table 2 Reasons for skipping prescribed dose(s) of medication

| Reason | Total n=321 |
|--------|-------------|
| I simply forgot | 43 |
| I was experiencing side effects | 22 |
| I was feeling better | 15 |
| Safety concerns | 13 |
| I did not think the treatment was working | 12 |
| Treatment did not make me feel better | 11 |
| I was no longer feeling better | 10 |
| I could no longer afford treatment | 8 |
| Reasons that were beyond my control | 11 |
| Other | 11 |
| Not sure/do not recall | 8 |

Notes: All patients who had taken a disease-modifying therapy prescription medication and skipped doses answered the following question: “Which of the following, if any, are reasons why you skip doses of your disease-modifying medication? Please select all that apply.”
Clear communication between patients and HCPs is key for optimal MS care.\textsuperscript{12,13} For patients, understanding what treatment will, and perhaps more importantly, will not do is important. Nearly one-third of patients who skipped doses did not tell their HCP, reinforcing the fact that patient adherence should be routinely evaluated. Forgetting was cited at a similar rate in the Global Adherence Project observational study (50%\textsuperscript{20} as in this survey (43%). Neurologists felt the main reason for non-adherence was side effects, indicating they are not aligned with patients on the main reasons for non-adherence. Identified treatment challenges also may affect adherence. For neurologists to effectively address non-adherence, they must understand patients’ reasons for non-adherence, dose skipping patterns, and treatment challenges for individual patients.

Limitations

The results of this study should be interpreted with caution, particularly concerning the representativeness of the unweighted populations (ie, European patients and neurologists). The European results are only representative of the patients and neurologists who completed the survey and cannot reliably be extrapolated to the entire MS population/treating neurologists. Although the translations were verified by native speakers, the multinational and multilingual nature of the survey may influence the results. Given the survey recruitment design and selection, response biases are possible and should be considered when interpreting the results. Patient recruitment targets were set for the total population per country without subtargets based on MS disease type; therefore, disease subtypes were not actively balanced across countries. This lack of balancing may be the reason for the unexpectedly high number of patients who identified as having PRMS in Germany compared with other countries or it may be an inconsistency in translation of the definition of PRMS. Also, we do not know which, if any, participating patients came from the same practices as the neurologists due to the independent recruitment of patients and neurologists. Therefore, the analysis represents both different perceptions and potentially different disease courses, treatment experiences, and patient–HCP interactions. Further, although the survey’s online approach allowed for a relatively large (900 neurologists and >900 patients) sample and an opportunity for respondents to answer sensitive questions honestly without introducing the bias of an interviewer, a bias associated with respondent self-selection may exist. We do not have data on the percentage of patients and neurologists who completed the survey compared with those who received the survey. The survey also is limited in that cognition was not assessed so, potentially, only patients with less cognitive impairment completed the survey. Finally, the differences in the importance and role of treatment cost between countries should be interpreted with caution, because the cause of these differences is likely due to the different health care systems. Along with costs, different health care systems also can bias perceptions with regard to treatment satisfaction and value of care.\textsuperscript{21}

Conclusion and recommendations

The results presented offer insight into differences between the perceptions of patients and neurologists and highlight differences in priorities for choosing a MS therapy. The study was aimed at identifying themes that might warrant further study with more robust methodology and was not in itself aiming to provide definitive statistics. Providing information that raises awareness of the importance of patient/neurologist collaboration and actions that can enhance these relationships should be considered. Therefore, a collaborative approach for setting appropriate treatment expectations and greater patient input in therapy choice is key for improving treatment satisfaction and adherence.

Based on these data, some action points were identified: 1) for HCPs, develop effective tools to improve communication, improving recognition of patients’ priorities, including patient-related outcomes; 2) for patients, develop educational programming that aids in learning about the disease and how to more actively participate in all aspects of decision making, thus working to become better self-advocates; and 3) opportunities for professional organizations (eg, European Academy of Neurology, American Academy of Neurology) and patient organizations (eg, European Multiple Sclerosis Platform) to work together to raise awareness of discrepancies in perceptions that exist between patients and HCPs.

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Author contributions
Mar Tintoré, Maggie Alexander, Martin Duddy, David E Jones, Nancy Law, Gilmore O’Neill, Antonio Uccelli, Robert Weissert, and Sibyl Wray participated in the State of MS Consortium, which designed the survey questions. Mar Tintoré, Maggie Alexander, Kathleen Costello, Martin Duddy, David E Jones, Nancy Law, Gilmore O’Neill, Antonio Uccelli, Robert Weissert, and Sibyl Wray interpreted and critically assessed the data for inclusion in the manuscript. Mar Tintoré, Maggie Alexander, Kathleen Costello, Martin Duddy, David E Jones, Nancy Law, Gilmore O’Neill, Antonio Uccelli, Robert Weissert, and Sibyl Wray contributed to and critically revised the manuscript.

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Table S1 Neurologist demographics

| Characteristics | Germany | Italy | Spain | UK | US | Total |
|-----------------|---------|-------|-------|----|----|-------|
| Neurologists (n) | 200     | 200   | 150   | 150| 200| 900   |
| Years in practice, n (%) | | | | | | |
| ≤10             | 80 (40) | 57 (29) | 50 (33) | 65 (43) | 45 (23) | 302 (34) |
| 11–20           | 78 (39) | 59 (30) | 67 (45) | 61 (41) | 63 (32) | 334 (37) |
| 21–30           | 36 (18) | 59 (30) | 26 (17) | 16 (11) | 49 (24) | 180 (20) |
| ≥31             | 6 (3)   | 25 (13) | 7 (5)  | 8 (5)  | 43 (22) | 85 (9)  |
| Mean (SD) age (years) | | | | | | |
| 194             | 174     | 170   | 235   | 124 | 180 | 180   |

Notes: Percentages may not add to 100 due to rounding. Weights were applied to the total data to give each country equal representation in the overall results. *Significantly different than Germany. †Significantly different than Italy. ‡Significantly different than Spain. §Significantly different than the UK. ¶Significantly different than the US.

Abbreviations: MS, multiple sclerosis; SD, standard deviation.

Table S2 Symptoms patients are not comfortable discussing with neurologist

| Symptoms (%) | Total patients (variable) | Total neurologists n=900 |
|--------------|---------------------------|------------------------|
| Sexual difficulties | 54                       | 87                     |
| Bladder/bowel problems | 28                      | 54                     |
| Mood swings    | 26                       | 37                     |
| Difficulty swallowing | 21                      | 4                      |
| Cognitive/memory issues/trouble concentrating | 21     | 37                     |
| Difficulty walking | 19                    | 3                      |
| Tremors        | 19                       | 2                      |
| Muscle spasms  | 18                       | 2                      |
| Pain           | 17                       | 4                      |
| Vision problems | 17                    | 3                      |
| Fatigue        | 16                       | 6                      |
| Dizziness/lightheadedness | 16            | 2                      |
| Speech problems | 16                    | 5                      |
| Difficulties with fine motor tasks | 15    | 5                      |
| Impaired balance/tendency to fall | 15        | 4                      |
| Tingling/“pins and needles” sensations, itching | 14    | 2                      |
| Numbness/impaired sensation | 13             | 2                      |
| Impaired upper limb function | 12        | 2                      |
| None           | N/A                      | 8                      |

Notes: Survey question: “Which of the following symptoms, if any, are you not comfortable talking about with your neurologist you see the most for your multiple sclerosis? Which of the following symptoms, if any, do you think your multiple sclerosis patients are not comfortable talking about with you? Please select all that apply.” Each symptom is based on those who experienced symptom (and also saw a health care provider in the last year).

Abbreviation: N/A, not applicable.
Figure S1 Impact of multiple sclerosis on the following aspects of your life/your patients’ lives.

Notes: Survey question: “How much of an impact, if any, has having multiple sclerosis had on each of the following aspects of your life/your patients’ lives?” Base: qualified patients who said the attribute was applicable; qualified neurologists who said they had an opinion on the attribute.

Table S3 Symptoms experienced

| Symptoms                                      | Have experienced | Experience daily |
|-----------------------------------------------|------------------|------------------|
|                                              | n=982            | (variable)       |
| Fatigue                                      | 71               | 74               |
| Tingling/“pins and needles” sensations, itching| 59               | 54               |
| Difficulty walking                           | 58               | 69               |
| Numbness/impaired sensation                  | 56               | 56               |
| Impaired balance/tendency to fall            | 56               | 57               |
| Pain                                         | 50               | 59               |
| Muscle spasms                                | 46               | 51               |
| Mood swings                                  | 45               | 46               |
| Vision problems                              | 41               | 44               |
| Impaired upper limb function                 | 40               | 46               |
| Bladder/bowel problems                       | 40               | 70               |
| Dizziness or lightheadedness                 | 39               | 38               |
| Cognitive/memory issues/trouble concentrating| 39               | 61               |
| Difficulties with fine motor tasks           | 33               | 53               |
| Sexual difficulties                          | 32               | 43               |
| Tremors                                      | 27               | 43               |
| Speech problems                              | 20               | 43               |
| Difficulty swallowing                       | 18               | 46               |
| Other                                        | 4                | 46               |

Notes: Base: all qualified patients who had experienced an applicable MS symptom. Survey question: “Which of the following symptoms of multiple sclerosis, if any, have you experienced? Please select all that apply.” Base: all qualified patients who had experienced an applicable MS symptom. Survey question: “Thinking of the symptoms you have experienced, which of the following, if any, do you experience on a daily basis? Please think only of the symptoms you experience when you are not having a relapse. Please select all those that apply.”

Abbreviation: MS, multiple sclerosis.
Figure S2 Patient participation in DMT decision making.

Notes: n=643 qualified patients with MS who had taken prescription medications responded to the question, “How much input did you have in choosing your current/past disease-modifying medication(s), compared to your health care provider?” Percentages (number) are shown. Caution: country-level results with small base size should be interpreted as directional in nature and, as such, frequencies are shown in addition to the percentages where applicable.

Abbreviations: DMT, disease-modifying therapy; HCP, health care provider; MS, multiple sclerosis.

Figure S3 Patients’ satisfaction with DMTs.

Notes: n=463 qualified patients with MS who were currently taking prescription medication responded to the question, “Overall, how satisfied are you with your current disease-modifying MS medication(s)?” Percentages (numbers) are shown. Caution: country-level results with small base size should be interpreted as directional in nature and, as such, frequencies are shown in addition to the percentages where applicable.

Abbreviations: DMT, disease-modifying therapy; MS, multiple sclerosis.
Figure S4 Top factors important for patients to know when making decisions about disease-modifying treatments for MS according to neurologists and patients.

Notes: n=982 qualified patients with MS responded to the question, “Which of the following, if any, are most important for you to know when making decisions about disease-modifying treatments?” n=900 qualified neurologists responded to the question, “Which of the following, if any, are the most important for your MS patients to know when making decisions about their disease-modifying treatments?” The top 8 of 20 factors selected by patients are shown. The percentage of respondents selecting each option is indicated as a data label for each bar.

Abbreviations: MRI, magnetic resonance imaging; MS, multiple sclerosis.