Marginal structural Cox models for estimating the association between beta-interferon exposure and disease progression in a multiple sclerosis cohort

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**WEB-APPENDIX**

**Web-Appendix 1: Rationale behind hypothesizing that cumulative relapses are lying on the causal path of β-IFN and disability progression**

The exact mechanism of action of the β-IFN drugs in MS has never been fully established and is one reason why estimating the effect of these drugs in MS is not straightforward. In the absence of randomization, establishing a causal link between drug exposure and outcome requires subject-specific knowledge and careful implementation of that knowledge in the analysis. Suggesting a plausible causal path is the first step.

Relapsing-remitting patients experience relapses followed by periods of remission in which partial or complete recovery occurs. Based on the results from randomized, double-blind, placebo-controlled studies, β-IFN treatments reduced the severity and frequency of relapses (1–5) and hence increased the period between relapses (4). Consequently, a patient has more time to recover from the residual disability left by the past relapse. This extended period of relapse-free time due to β-IFN exposure may eventually contribute to a slower progression of disability (2–4). However, it should be noted that while most natural history studies indicate that long-term there is minimal or no association between relapse rates and disability progression, a specific window of opportunity for relapses to contribute to disease progression may exist (6, 7).

Therefore, we hypothesized that within a short time interval the cumulative relapses are acting as an intermediate variable for the treatment and disability progression relationship, i.e., the relapse frequency is influenced by prior β-IFN treatment and a greater (lesser) relapse frequency will result in faster (slower) disability progression. Also, we assume that the cumulative relapse count in the previous time period is a confounder that may dictate the treatment choice in subsequent time periods. Furthermore, experiencing an increased number of cumulative relapses after initiating treatment will increase the probability of discontinuing treatment (8). Hence, in this relationship, cumulative relapse is treated both as an intermediate variable and a confounder.

The causal path described above could be considered as rather simplistic. It is possible that cumulative relapse and disability progression have an unmeasured common cause (for example, low serum vitamin D levels). Should this data be available, then we would add that variable to the causal path between cumulative relapse and EDSS. Cumulative relapse would still be a time-dependent confounder and would need to be adjusted for accordingly.
Web-Appendix 2: Rationale behind using a marginal structural Cox model (MSCM) instead of a Cox model

For a longitudinal study with $N$ patients, let $i = 1, 2, \ldots, N$ be the patient index, $t = 0, 1, \ldots, T_i$ months be the follow-up time index, $A_i$ be the binary treatment status at month $t$ ($1 =$ treated, $0 =$ untreated), and $L_{it}$ be the baseline covariates of patient $i$. One possible model would express the hazard function of the time-dependent Cox model as follows:

$$\lambda_i(t | L_{i0}) = \lambda_{0t} \exp \left( \beta_1 A_{it} + \beta_2 L_{i0} \right), \quad (1)$$

where $\lambda_{0t}$ is the unspecified baseline hazard function, $\beta_2$ is the vector of log hazard ratios (HRs) for the baseline covariates and $\beta_1$ is the log HR of the current $\beta$-IFN status ($A_{it}$).

Assuming no tied event times, we estimate $\beta = (\beta_1, \beta_2)$ by maximizing the partial likelihood (9):

$$PL(\beta) = \prod_{i=1}^{N} \prod_{t=0}^{T_i} \left( \frac{Y_{it} \exp \left( \beta_1 A_{it} + \beta_2 L_{i0} \right)}{\sum_{k=1}^{N} Y_{kt} \exp \left( \beta_1 A_{kt} + \beta_2 L_{k0} \right)} \right)^{dN_{it}},$$

where $Y_i$ denotes whether patient $i$ belongs to the risk set at time $t$, $N_i$ is the number of events in the interval $[0, t]$ and $dN_{it}$ denotes the number of new events for patient $i$ at month $t$ (increment from month $t-1$, if any). This setting is more general than our case, where $N_i \leq 1$ and $dN_{it} = 1$ for at most 1 month.

However, ignoring the time-dependent confounder $L_{it}$ (i.e., an intermediate variable lying in the causal pathway of the treatment and the outcome) may lead to a biased estimate of $\beta$. Simply including this variable in the Cox model as a covariate as,

$$\lambda_i(t | L_{i0}, L_{it}) = \lambda_{0t} \exp \left( \beta_1 A_{it} + \beta_2 L_{i0} + \beta_3 L_{it} \right), \quad (2)$$

may still produce a biased estimate if $L_{it}$ is influenced by past exposure (10).

Inverse probability of treatment and censoring weights (IPTC; say $w, sw, w_{sw}, sw_{sw}$) are person-time specific measures of the degree to which a time-dependent variable confounds the treatment selection and censoring processes. These are used in the time-dependent Cox model to weight the contribution of each person-time observation so that confounding due to $L_{it}$ is
removed without changing the target parameter. In this way, MSCM facilitates correction for
time-dependent confounding. In the MSCM, these IPTC weights are inserted in the partial
likelihood function as follows (11–13):

$$PL_w(\beta) = \prod_{i=1}^{N} \prod_{t=1}^{T_i} \left( \frac{Y_{it} \exp(\beta_1 A_{it} + \beta_2 L_{it0})}{\sum_{k=1}^{N} Y_{kt} w_{kt} \exp(\beta_1 A_{kt} + \beta_2 L_{k0})} \right)^{dN_i \cdot w_i}.$$ 

The gradient with respect to the parameter vector $\beta$ of the log of the weighted partial likelihood
$PL_r(\beta)$ yields the score function $U_r(\beta)$. Equating $U_r(\beta)$ to zero yields a set of estimating equations
that can be solved using an iterative method such as the Newton-Raphson algorithm or a
penalized partial likelihood approach.
Web-Appendix 3: Approximation of the marginal structural Cox model

Let $D_t$ be an indicator of reaching EDSS 6 for the first time between the months $t-1$ and $t$. The data for patients who did not reach sustained EDSS 6 and remained uncensored until follow-up month $t$ can be modelled using the pooled logistic regression (logistic regression pooled over persons and times):

\[
\text{logit}[Pr(D_t = 1|D_{t(t-1)} = 0, A_t, L_{t0})] = \gamma_0(t) + \gamma_1A_t + \gamma_2L_{t0}.
\] (3)

Here $\gamma(t)$ is a smooth function of the month index $t$, represented as a restricted cubic spline, which is often used to reduce weight variability. Just as for cubic polynomial regression, use of a restricted cubic spline forces the relationship to be smooth even on the edges (14, chapter 6); see the R code in the Web-Appendix 5. The log OR of the current $\beta$-IFN status in this pooled logistic regression, $\gamma_1$, is generally a good approximation of the corresponding log hazard ratio obtained from the time-dependent Cox model ($\beta_1$), provided that censoring is ignorable (15) and relatively short intervals are chosen so that the probability of outcome occurrence in each time interval is small (16, 17). The corresponding likelihood function can be expressed as:

\[
L(\gamma) = \prod_{i=1}^N \prod_{t=0}^{T_i} p_{it}^{D_{it}} (1 - p_{it})^{(1-D_{it})},
\]

where $\gamma = (\gamma_0, \gamma_1, \gamma_2)$ and $\text{logit}(p) = \gamma_0(t) + \gamma_1A_t + \gamma_2L_{t0}$.

Hernán et al. (10) suggested use of weighted pooled logistic regression to approximate MSCM (IPTC weighted time-dependent Cox model) estimates of treatment association ($\beta_1$) and others have followed this suggestion. (18–22). The weighted likelihood function is then written as (15):

\[
L_w(\gamma) = \prod_{i=1}^N \prod_{t=0}^{T_i} \left(p_{it}^{D_{it}} (1 - p_{it})^{(1-D_{it})}\right)^{w_i}.
\]

This approximate approach was suggested mainly because software available at that time was unable to handle patient-specific time-varying weights in a Cox model. It has been noted that this approximation approach is inadequate when the event is not rare (23). Subsequently Xiao et al. (24) suggested the direct use of the Cox model weighted by IPTC weights to overcome this limitation. Through simulation, these authors also showed that direct use of the Cox model weighted by IPTC weights instead of any approximate MSCM approach (10) considerably reduced the variability of the estimated treatment association, even when both methods use the same weights.
Web-Appendix 4: Weight models

The stabilized IPT weights for patient $i$ at month $t$ are expressed as:

$$sw_{it}^T = \prod_{j=0}^{t} \frac{pr(A_{ij} = a_j | A_{i(j-1)}, L_{i0} = l_{i0})}{pr(A_{ij} = a_j | \bar{A}_{i(j-1)}, \bar{L}_{i0} = \bar{l}_{i0})}.$$  (4)

The probability appearing in the numerator of $sw^T$ is modeled using a pooled logistic model as follows:

$$logit[pr(A_{ij} | A_{i(j-1)}, L_{i0})] = \alpha_0(j) + \alpha_1 A_{i(j-1)} + \alpha_2 L_{i0},$$  (5)

where treatment status at the previous time interval ($A_{ij-1}; A_i = 0$ for all patients), the baseline covariates ($L_0$; in our application, EDSS, age, disease duration, sex) and a restricted cubic spline of the follow-up month index are included as predictors. These covariates, as well as the time-varying confounder cumulative relapse ($L_m$) and its interaction with prior treatment status are included in the denominator model:

$$logit[pr(A_{ij} | \bar{A}_{i(j-1)}, \bar{L}_{i0}, \bar{L}_j)] = \alpha_0(j) + \alpha_1 A_{i(j-1)} + \alpha_2 L_{i0} + \alpha_3 L_{j0} + \alpha_4 L_{j0} + \alpha_5 A_{i(j-1)}L_j.$$  (6)
The output of this fit is reported in Web-Table 1.

**Web-Table 1**: Estimated coefficients from the treatment model (denominator of \(sw_{jt}\)) for patients with relapsing-onset multiple sclerosis (MS), British Columbia, Canada (1995-2008)

<table>
<thead>
<tr>
<th></th>
<th>Estimate</th>
<th>z-value</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\beta)-IFN(_j)</td>
<td>9.78</td>
<td>102.92</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>EDSS(^a)</td>
<td>0.12</td>
<td>4.31</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Age(^{a,b})</td>
<td>-0.07</td>
<td>-1.70</td>
<td>0.09</td>
</tr>
<tr>
<td>Disease duration(^{a,b})</td>
<td>-0.17</td>
<td>-3.17</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Sex(^a)</td>
<td>-0.07</td>
<td>-0.96</td>
<td>0.34</td>
</tr>
<tr>
<td>Cumulative relapse</td>
<td>0.34</td>
<td>7.70</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Cumulative relapse;(\beta)-IFN(_j)</td>
<td>-0.55</td>
<td>-10.83</td>
<td>&lt; 0.001</td>
</tr>
</tbody>
</table>

EDSS, expanded disability status scale.

\(^a\) Time index is also fitted with restricted cubic spline, but the corresponding coefficients are not reported in the table.

\(^a\) Baseline covariates (\(L_0\)).

\(^b\) Expressed in decades.

The predicted value from the (denominator) model (.6) yields the estimated probability of the patient’s treatment status in that month \(t\). Since the exposure status may vary from one time point to another, first we estimate the probability of the observed treatment status at each time point, and then obtain the probability of the observed exposure sequence of a given patient by multiplying the corresponding probabilities. The numerator of \(sw_{jt}\) is estimated in a similar fashion from model (.5), where \(L_{ij}\) is not included as a predictor. Dividing the numerator model probabilities of the patient’s observed treatment status \(a_{ij}\) (either 0 or 1) by the corresponding denominator model probabilities yields the estimated IPT weights \(sw_{jt}\) that account for the confounding due to \(L_{ij}\), given the required assumptions are met.

To estimate the IPTC weights \(sw_{jt} = sw_{jt} \times sw_{jc}\), the inverse probability of censoring (IPC) weights \(sw_{jc}\) are estimated in the same fashion. In order to produce the normalized IPTC weights \(sw_{n}\), each weight \(sw\) is divided by its risk set’s mean weight.
Web-Appendix 5: MSCM fitting in R

For time-dependent survival analysis, all person-time observations are pooled to make an augmented dataset. Short intervals, such as months, are chosen so that the most recently observed changes of the time-varying variables can be updated in a new row in the dataset to reflect the patient’s time-varying status with respect to covariates, censoring and response. In the longitudinal analysis literature, this is referred to as the ‘long’ format.

Guidelines regarding IPTC weight calculations in R are available in the literature (21). These IPTC weights can be viewed as a generalization of the Horvitz-Thompson estimator (25–27). Recently, due to the availability of packages for the analysis of complex surveys in standard software (SAS, Stata and R), it is possible to fit the time-dependent IPTC weighted Cox model directly or via approximation, say, using the weighted pooled logistic model. In all the model choices, reliable SEs can be obtained from a reasonable number of patient-specific bootstrap samples.

- Most MSCM analyses in the literature use weighted pooled logistic regression to approximate the IPTC weighted Cox model fit. In R, performing weighted pooled logistic regression using the glm function from the base package (with log link) is straightforward (21).
- Similarly, the svyglm function from the survey package can be used to implement the (weighted) pooled logistic model (26).
- With data organized in person-month format, to perform survival analysis using the weighted Cox model, we used the Andersen-Gill’s counting process approach as implemented in the svycoxph function from the R package survey (28) with the weights option. Approximation via complementary-log-log and Poisson models can also be implemented using the same package. A sample code follows:

```r
require(survey)
require(rms)
(weighted.design<-svydesign(id=~ID, data=long.format, 
                           weight=~normalized.stabilized.weight))
svycoxph(Surv(start, stop, event) ~ drug + covariate.list, 
         design=weighted.design)
svyglm(event ~ drug + rcs(Time) + covariate.list, 
       family=binomial(link=log), design=weighted.design)
svyglm(event ~ drug + rcs(Time) + covariate.list, 
       family=binomial(link=cloglog), design=weighted.design)
svyglm(event ~ offset(log(stop-start)) + drug + rcs(Time) + 
       covariate.list, family=poisson(), design=weighted.design)
```
Alternatively, the `coxph` function from the `survival` package (29) can be used to fit the weighted Cox model (24). To handle correlated observations, the `cluster` option must be specified to identify the person-month observations from the same patient. Robust SEs are obtained by specifying the option `robust = TRUE`. 
Web-Appendix 6: Exclusion criteria and summary of selected cohorts

In total, 2,671 patients met the eligibility criteria to receive β-IFN treatment between July 1995 and December 2004 (31). Of these, patients who were exposed to a non–β-IFN immunomodulatory drug, a cytotoxic immunosuppressant for MS (n=172), or an MS clinical trial (n=21) prior to baseline were excluded from the analysis. If the exposure occurred after baseline, data were censored at the start of the exposure of the non-β-IFN treatment. Other exclusion criteria included unknown MS onset date (n=10), insufficient EDSS measurements (n=436), reaching the outcome (n=218) or the secondary progressive stage before the eligibility date (n=217). Some patients met multiple exclusion criteria.

As a result, 1,697 patients were selected. A summary of their characteristics are reported in Web-Table 6.

Web-Table 2: Characteristics of the selected cohort of patients with relapsing-onset multiple sclerosis (MS), British Columbia, Canada (1995-2008).

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>β-IFN exposed patients</th>
<th>β-IFN unexposed patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frequency</td>
<td>868</td>
<td>829</td>
</tr>
<tr>
<td>Women, n (%)</td>
<td>660 (76.0)</td>
<td>637 (76.8)</td>
</tr>
<tr>
<td>Disease duration (at baseline)</td>
<td>5.8± (6.6σ)</td>
<td>8.3± (8.5σ)</td>
</tr>
<tr>
<td>Age (at baseline)</td>
<td>38.1± (9.2σ)</td>
<td>41.3± (10.0σ)</td>
</tr>
<tr>
<td>EDSS score (at baseline)</td>
<td>2.0± (0-6.5σ)</td>
<td>2.0± (0-6.5σ)</td>
</tr>
<tr>
<td>Relapse rate / year (over the 2 years prior to baseline)</td>
<td>0.5± (0-1.2σ)</td>
<td>0.5± (0-1.0σ)</td>
</tr>
<tr>
<td>Person-years exposed to β-IFN treatment</td>
<td>2,530</td>
<td>0</td>
</tr>
<tr>
<td>Person-years not exposed to β-IFN treatment</td>
<td>1,400</td>
<td>2,960</td>
</tr>
</tbody>
</table>

EDSS, expanded disability status scale.

a Mean  
b Standard Deviation  
c Median  
d Range  
e IQR
Web-Appendix 7: Sensitivity analysis: impact of weight trimming

If the weights contain extreme values, one should be concerned about the positivity assumption. The MSCM approach is built on the counterfactual framework and it is necessary to assume patients could choose treatment exposure or non-exposure at any time point. If a group of patients with similar covariate history rarely or never receive treatment, then the estimated probability of being treated would be close to zero. Conversely, if a group of patients with similar covariate history almost always or always receive treatment, then the estimated probability of being treated would be close to one. Then the corresponding fitted probability will be close to zero or one resulting in a very large or small inverse probability weight respectively. This may produce unstable estimates from the MSCM.

As a sensitivity analysis, one could restrict the analysis to the subset of patients that have a probability of treatment and censoring that is reasonably removed from 0 and 1 at every time point. This procedure is known as trimming (30). As with truncation of the weights, systematically excluding such patients may produce a biased estimate. Also, the interpretation may lack generalizability due to this restriction. However, since the patients with extreme weights are removed, a relatively stable point estimate with a smaller CI would be expected.

After estimating the fitted probabilities from the weight models, if the probabilities are such that a few person-time observations are contributing too much in the pseudo-population, this may make the estimate of the causal association unstable. In our sensitivity analysis, we removed the patients with at least one fitted value either greater than 0.95 or less than 0.05 (represented more than 20 times in the pseudo-population). This left 1,603 patients, with 133 reaching the outcome. MSCM using swon led to a HR estimate of 1.33 with a 95% bootstrap CI of 0.94 - 1.89. The conclusion regarding the treatment association between β-IFN and time to sustained EDSS 6 from these results remained the same.
Web-Appendix 8: Sensitivity analysis: impact of more restrictive eligibility criteria

As another sensitivity analysis, a more restricted study sample was selected by defining active disease (two or more documented relapses during the two years prior to baseline) as part of the eligibility criteria, while also including all the previous criteria. This left 747 patients in the study with 3028 person-years of follow-up and 1460 person-years of β-IFN exposure. Only 52 of these patients reached the irreversible disease outcome.

The model fit is reported in Web-Table 2. The regression coefficients and HR estimates were qualitatively similar to those reported in Table 2. The CIs from this restricted dataset were wider due to the smaller sample size. Still, the conclusion regarding the treatment association between β-IFN and time to sustained EDSS 6 remained the same as before.

Web-Table 3: The marginal structural Cox model (MSCM) fit with the normalized stabilized IPTC weights \(sw^{\text{net}}\) for time to sustained EDSS 6 to estimate the causal association between β-IFN treatment for patients with relapsing-onset multiple sclerosis (MS), British Columbia, Canada (1995-2008) selected by more restrictive eligibility criteria. The model was also adjusted for baseline covariates EDSS, age, disease duration and sex.

<table>
<thead>
<tr>
<th>Covariate</th>
<th>Estimate(^a)</th>
<th>HR  (^b)</th>
<th>95% CI (^c)</th>
</tr>
</thead>
<tbody>
<tr>
<td>β-IFN</td>
<td>0.18</td>
<td>1.19</td>
<td>0.68 - 2.11</td>
</tr>
<tr>
<td>EDSS</td>
<td>0.40</td>
<td>1.48</td>
<td>1.24 - 1.77</td>
</tr>
<tr>
<td>Disease duration(^e)</td>
<td>-0.14</td>
<td>0.87</td>
<td>0.55 - 1.37</td>
</tr>
<tr>
<td>Age(^e)</td>
<td>0.45</td>
<td>1.57</td>
<td>1.14 - 2.18</td>
</tr>
<tr>
<td>Sex(^f)</td>
<td>-0.33</td>
<td>0.72</td>
<td>0.38 - 1.35</td>
</tr>
</tbody>
</table>

HR, Hazard ratio; CI, confidence interval; EDSS, expanded disability status scale.

\(^a\) Estimated log HR: negative value is indicative of a beneficial association and positive value is indicative of a harmful association.

\(^b\) HR, indicating the instantaneous risk of reaching sustained and confirmed EDSS 6.

\(^c\) Based on 500 nonparametric bootstrap sample estimates.

\(^d\) 95% CI that does not include 1.

\(^e\) Expressed in decades.

\(^f\) Reference level: Male.
Web-Appendix 9: Sensitivity analysis: impact of the cumulative exposure to β-IFN

We also assessed the impact of the cumulative exposure to β-IFN (proportion of months exposed) over the last two years on time to sustained EDSS 6. The model fit is reported in Web-Table 3. This analysis also failed to detect a significant association between the cumulative exposure to β-IFN and the hazard of reaching sustained EDSS 6. A similar finding was observed when the cumulative exposure was restricted to the past year only (data not shown).

**Web-Table 4:** The marginal structural Cox model (MSCM) fit with the normalized stabilized IPTC weights \( w^{ou} \) for time to sustained EDSS 6 to estimate the causal association of cumulative exposure to β-IFN over the last two years for patients with relapsing-onset multiple sclerosis (MS), British Columbia, Canada (1995-2008). The model was also adjusted for baseline covariates EDSS, age, disease duration and sex.

<table>
<thead>
<tr>
<th>Covariate</th>
<th>Estimate</th>
<th>HR (^b)</th>
<th>95% CI (^c)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cumulative β-IFN(^a)</td>
<td>0.53</td>
<td>1.70</td>
<td>0.64 - 4.53</td>
</tr>
<tr>
<td>EDSS</td>
<td>0.54</td>
<td>1.71</td>
<td>1.53 - 1.91(^d)</td>
</tr>
<tr>
<td>Disease duration(^e)</td>
<td>-0.20</td>
<td>0.82</td>
<td>0.66 - 1.10</td>
</tr>
<tr>
<td>Age(^e)</td>
<td>0.30</td>
<td>1.34</td>
<td>1.10 - 1.63(^d)</td>
</tr>
<tr>
<td>Sex(^f)</td>
<td>-0.23</td>
<td>0.79</td>
<td>0.55 - 1.15</td>
</tr>
</tbody>
</table>

HR, Hazard ratio; CI, confidence interval; EDSS, expanded disability status scale.

\(^a\) Expressed as proportion of months exposed over last two years.
\(^b\) HR, indicating the instantaneous risk of reaching sustained and confirmed EDSS 6.
\(^c\) Based on 500 nonparametric bootstrap sample estimates.
\(^d\) 95% CI that does not include 1.
\(^e\) Expressed in decades.
\(^f\) Reference level: Male.
Web-Appendix 10: Sensitivity analysis: impact of the cumulative number of relapses in the last year

We also assessed the impact of the exposure to \(\beta\)-IFN on time to sustained EDSS 6 while considering the cumulative number of relapses in the last year (instead of the last two years) as the time-varying confounder. The model fit is reported in Web-Table 5. This analysis also failed to detect a significant association between the exposure to \(\beta\)-IFN and the hazard of reaching sustained EDSS 6.

**Web-Table 5:** The marginal structural Cox model (MSCM) fit with the normalized stabilized IPTC weights \(sw^{(n)}\) for time to sustained EDSS 6 to estimate the causal association of exposure to \(\beta\)-IFN for patients with relapsing-onset multiple sclerosis (MS), British Columbia, Canada (1995-2008) while considering the cumulative number of relapses in the last year as the time-varying confounder. The model was also adjusted for baseline covariates EDSS, age, disease duration and sex.

<table>
<thead>
<tr>
<th>Covariate</th>
<th>Estimate</th>
<th>HR (^a)</th>
<th>95% CI (^b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\beta)-IFN</td>
<td>0.31</td>
<td>1.36</td>
<td>0.96 - 1.92</td>
</tr>
<tr>
<td>EDSS</td>
<td>0.54</td>
<td>1.72</td>
<td>1.54 - 1.92</td>
</tr>
<tr>
<td>Disease duration(^d)</td>
<td>-0.18</td>
<td>0.82</td>
<td>0.66 - 1.04</td>
</tr>
<tr>
<td>Age(^d)</td>
<td>0.28</td>
<td>1.32</td>
<td>1.10 - 1.60</td>
</tr>
<tr>
<td>Sex(^e)</td>
<td>-0.22</td>
<td>0.80</td>
<td>0.55 - 1.16</td>
</tr>
</tbody>
</table>

HR, Hazard ratio; CI, confidence interval; EDSS, expanded disability status scale.

\(^a\) HR, indicating the instantaneous risk of reaching sustained and confirmed EDSS 6.

\(^b\) Based on 500 nonparametric bootstrap sample estimates.

\(^c\) 95% CI that does not include 1.

\(^d\) Expressed in decades.

\(^e\) Reference level: Male.
WEB-REFERENCES


