The Effect of Body Mass Index on Brain Volume and Cognitive Function in Relapsing–Remitting Multiple Sclerosis: A CombiRx Secondary Analysis

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Introduction
Multiple sclerosis (MS) is a neurological autoimmune, inflammatory degenerative disease leading to physical, emotional, and cognitive disability among young adults.1-2 Aside from race and genetics, which are non-modifiable risk factors for MS, recent studies have shown an association between risk of MS in adolescents and young adults and modifiable factors, such as obesity.3 In fact, obesity has been recognized recently as a modifiable emerging risk factor for MS by the American Academy of Neurology (AAN),4 with 70% of MS patients obese or overweight.5,6

Obesity has been linked with cognitive dysfunction and brain volume loss in healthy adults,7,8 and brain volume loss has been recognized as one of the best predictors for cognitive impairment in MS.9,10 Although multiple studies have shown a link between cognitive impairment and brain volume loss in MS, it is unclear if this association is triggered by modifiable risk factors such as body mass index (BMI) and/or non-modifiable factors, such as genetics, or a combination of both. Cross-sectional studies,11-18 provided preliminary and partial support for the relationship between high BMI and brain volume loss and cognitive dysfunction in MS, but there are also conflicting results. For example, Bove and colleagues (2019) and Galioto and colleagues (2019) did not show an association between BMI and cognition but Owji and colleagues (2019) demonstrated a negative correlation between BMI and cognitive function as measured by the Paced Auditory Serial Addition Test (PASAT) and Symbol Digit Modalities Test (SDMT). The role of BMI in MS continues to be controversial; therefore, there is a

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ABSTRACT

BACKGROUND: Multiple sclerosis (MS) is an autoimmune disease leading to physical, emotional, and cognitive disability. High body mass index (BMI) may impact cognitive function and brain volume in MS. Yet, there is paucity of evidence addressing the impact of BMI on cognitive function and brain volume in MS.

OBJECTIVES: The purpose of this study was to examine the effects of BMI on normal appearing brain volume and cognitive function in patients with relapsing–remitting MS.

METHODS: A secondary data analysis of the NIH CombiRx study was conducted. Multivariate regression and mixed model analyses were executed to analyze the effect of BMI on brain volume and cognitive function.

RESULTS: The mean baseline age of the 768 participants was 38.2 (SD = 9.4) years. 73% were female and 88.8% were Caucasian. The mean BMI was 28.8 kg/m2 (SD = 6.7). The multivariate regression and mixed model analyses failed to show a clinical effect of BMI on brain volume and cognitive function.

CONCLUSION: BMI did not show an effect on cognitive function and brain volume among MS patients. Although there is increased interest in the effects of modifiable factors on the course of MS, the effects of BMI on brain volume and cognitive function are debatable and warrant further research.

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KEYWORDS: Multiple sclerosis, RRMS, MRI, cognition, brain volume, outcome measurements

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critical need for gaining a better understanding of the effect of BMI on cognitive function and brain volume in this population.

This study addressed a major methodological limitation of prior studies. In particular, most of the prior studies have been cross-sectional studies as compared to this study, which was based on a longitudinal randomized control trial (RCT) for 3 years. The purpose of this study was to examine the effects of BMI on normal appearing brain volume and cognitive function in adult patients with relapsing-remitting MS (RRMS) treated with interferon-β or glatiramer acetate while controlling for potential confounders of age, sex, ethnicity, duration of illness from diagnosis and from first symptom, relapses, disability, MS medications, and smoking. The revised Scaffolding Theory of Aging and Cognition (STAC-R) guided this study. The STAC-R consists of a model linking lifestyle activities, biological factors, cognition, and brain volume, depicting life course experiences that may enrich or deplete neuronal functions. The scaffolding model suggests that individuals with MS who accrue multiple neural insults throughout the course of their illness will exhibit loss of brain volume and poor cognitive function. Hence, we hypothesized that high BMI may accelerate brain volume loss and cognitive dysfunction.

Methodology

Parent Trial—CombiRx and ethics statement

This study is a secondary data analysis of the CombiRx trial, a phase III, multicenter RCT sponsored by the National Institutes of Neurological Disorders and Stroke (US NIH Grant/Contract U01NS045719, R21NS41986; NIH identifier number NCT00211887). This trial randomized individuals to one of three disease-modifying therapies (DMTs): interferon-β (25%), glatiramer acetate (25%) or both interferon and glatiramer acetate (50%). Participants were naïve to treatment at entry. The CombiRx trial was approved by the applicable central or institutional review boards and the Data and Safety Monitoring Committee (DSMC) appointed by National Institutes of Health (NIH)/National Institute of Neurological Disorders and Stroke (NINDS) before site initiation and recruitment of participants. Written, informed consent was obtained prior to any screening procedures or enrollment. The trial was registered at www.clinicaltrials.gov/ct2/show/NCT00211887; for more information, please see www.CombiRx.org). The present study utilized deidentified data for the secondary analysis and was considered exempt by Mount Sinai Icahn School of Medicine institutional review board (IRB-16-1247) and New York University institutional review board (IRB-FY2019-2463).

Secondary Analysis Study

A total of 768/1008 patients completed the 3 years and were included in this secondary analysis, as they had sequential brain MRI films and brain volume calculations, BMI, and cognitive function measurements for 3 years. White and gray matter volume at 36th month was missing in 219 patients with RRMS in the CombiRx study (21.7%). Intention to treat (ITT) analysis of the full sample (n = 1008) was performed with 5 replicates of imputation based on BMI and brain volume using IBM SPSS version 23. There were no exclusion criteria in this secondary data analysis.

Study Measures

BMI was calculated based on weight in kilograms divided by the square of height in meters, which were measured during the study every 3 months by the research team. BMI was evaluated both as a continuous variable and as a categorical variable divided into normal (<25 kg/m²), overweight (25-29.9 kg/m²), and obese (≥30 kg/m²). Brain volume was acquired using a standardized protocol that included 7 separate scan series. The brain volume analyzed in this study was based on the normal appearing white matter (NAWM) and normal appearing gray matter (NAGM). MRI abnormalities are seen in NAWM and NAGM in early RRMS and the lack of correlation between NAWM or NAGM and lesion abnormalities suggests that they are developed by partly independent mechanisms. Cognitive function was assessed by the PASAT. The PASAT tests memory, speed of information processing, concentration, and attention, with scores from 0–60 while higher scores indicating better cognitive performance. In MS studies, the PASAT Cronbach’s alpha was .90 and the test–retest coefficients ranged between .90 and .97.

Assessment of construct validity of the PASAT showed good correlations with other cognitive tests on attention, working memory, processing ability, and speed. The CombiRx study included 2 PASAT screening visits before the baseline visit in an effort to diminish the well-known learning curve for the test. Substantial improvement of the PASAT between the first and second screening visits was observed, with a smaller change between the second screening visit and baseline. Although the assumption of homoscedasticity of errors of the PASAT was met, it had multiple outliers and its residuals did not follow a normal distribution. Therefore, in this study, the PASAT was dichotomized with a cutoff of 53 based on the PASAT median of the sample. Continuous independent variables included age, duration of illness from first symptom in years, and number of relapses in the last 3 years. Categorical variables included sex, race (Caucasian, African American, other), disease modifying therapies (DMTs), systolic and diastolic blood pressure, and smoking (ex-smoker, never smoked and current smoker).

Lastly, disability was measured by the Expanded Disability Status Scale (EDSS) with entry criteria of EDSS ≤6.0. The upper limit of the EDSS was determined by the CombiRx researchers. Their goal was to include only naïve relapsing-remitting patients and not those that transition into secondary progressive MS. The EDSS score was determined.
based on a neurological exam every 3 months by the neurologist. The EDSS quantifies disability in MS with scores from 0–10 in .5 point increments with higher scores indicating higher disability and monitors changes in the level of disability over time.

**Data Analysis and Management**

Excel data sheets with de-identified data provided by the principal investigators of the CombiRx trial were imported into IBM SPSS version 23 for data analyses. The statistical significance for this secondary data analysis study with non-directional hypotheses was set at a 2-tailed alpha level of .05. Descriptive statistics were performed to portray the sample characteristics. In addition, Chi-square and t-tests were performed to analyze the association between categorical and continuous variables. Multivariate linear regression, multivariate logistic regression, and mixed model of interaction with time analyses were executed to analyze the effect of BMI on brain volume or cognitive function while controlling for age, sex, ethnicity, smoking, blood pressure, disease duration, relapses, and disability.

**Results**

**Demographics and Disease Related Variables**

The mean baseline age of the 768 participants was 38.2 (SD = 9.4) years, ranging from 18 to 60 years, and a median age of 38 years. Seventy-three percent were female and 88.8% were Caucasian. The mean duration of illness was 4.2 years based on time from first symptom (Table 1; Table 2). The baseline mean BMI was 28.8 (SD = 6.7) kg/m² ranging from 16 kg/m² to 60 kg/m². A total of 32.7% had a normal BMI (≤24.99 kg/m²), 31% were overweight (25-29.99 kg/m²), and 36.3% were obese (≥30 kg/m²). A test for trend did not demonstrate a meaningful change in BMI during the course of the study (Table 3). The mean baseline PASAT was 50.02 (SD = 10.40) and the 36th month PASAT was 53.84 (SD = 8.36). There was no significant correlation between the baseline BMI and the 36th month PASAT scores (Pearson r = .03, spearman rho = .05 p’s > .05).

| CHARACTERISTICS | (N) (%) |
|-----------------|--------|
| **Sex**         |        |
| Male            | 207    | 27    |
| Female          | 561    | 73    |
| **Race**        |        |
| Caucasian       | 682    | 88.8  |
| African american| 52     | 6.8   |
| Other           | 34     | 4.4   |
| **Baseline age**|        |
| ≤29             | 160    | 20.8  |
| 30–39           | 265    | 34.5  |
| 40–49           | 241    | 31.4  |
| ≥50             | 102    | 13.3  |
| **Marital status**|      |
| Married         | 473    | 61.6  |
| Single          | 220    | 28.6  |
| Divorced        | 61     | 7.9   |
| Separated       | 14     | 1.8   |
| **Smoking history**|    |
| Ex-smoker       | 166    | 24.1  |
| Never smoker    | 348    | 50.6  |
| Current smoker  | 174    | 25.3  |
| **Family history of MS**|   |
| No              | 595    | 77.5  |
| Yes             | 173    | 22.5  |
| **MS medications**|   |
| Glatiramer acetate| 212  | 27.6  |
| Interferon      | 178    | 23.2  |
| Interferon+Glatiramer acetate| 378  | 49.2  |
| **Baseline BMI (3 categories)**| |
| Normal BMI      | 251    | 32.7  |
| Overweight      | 238    | 31    |
| Obese           | 279    | 36.3  |
| **BMI at 36th month (3 categories)**| |
| Normal BMI      | 232    | 30.2  |
| Overweight      | 246    | 32    |
| Obese           | 290    | 37.8  |

**Table 1.** Sample characteristics: Demographic and clinical categorical characteristics of the study participants (n = 768).

| CHARACTERISTICS | MEAN | SD  |
|-----------------|------|-----|
| Age in years    |      |     |
| Baseline age    | 38.2 | 9.4 |
| Disease duration in years | | |
| From 1st symptom| 4.2  | 5.2 |
| Relapses        |      |     |
| Number of relapses in last 3 years | 2.4  | 0.9 |
| Disability (EDSS)|   |
| At baseline     | 1.9  | 1.1 |
| At 36th month   | 1.9  | 1.3 |
| BMI             |      |     |
| Baseline BMI    | 28.81| 6.74|
| BMI at 36th month| 29.04| 6.64|
| NAWM            |      |     |
| Baseline        | 469.22| 54.51|
| At 36th month   | 467.68| 58.29|
| NAGM            |      |     |
| Baseline        | 588.07| 63.42|
| At 36th month   | 584.60| 61.51|

**Table 2.** Sample characteristics: Demographic and clinical continuous characteristics of the study participants (n = 768).
The MS patients’ cognitive function has improved over the 3 years of the study ($X^2(1) = 113.64, P < .001$). Explicitly, the PASAT scores improved from baseline to month 12 and month 24 but were stable between month 24 and month 36. The analysis showed that patients missing the PASAT at month 36th were those with lower baseline PASAT scores. DMT assignment did not significantly predict improvement in PASAT scores.

**BMI, Cognitive Function, NAWM, and NAGM Brain Volume**

The multivariable logistic regression analyses and the mixed model analysis failed to show an effect of BMI on cognitive function in patients with RRMS. BMI was evaluated for effects on cognitive function as a continuous variable and as a categorical variable due to its meaningful clinical implications. Least squares means adjusted for age, sex, race, and treatment group showed mean values of 2.7 for BMI < 25 kg/m$^2$; 3.2 for those ≥25 kg/m$^2$ and <30 kg/m$^2$; and 2.0 for those with BMI ≥30 kg/m$^2$ ($P = .21$).

BMI was evaluated for effects on normal appearing brain volume as a continuous variable and as a categorical variable due to its meaningful clinical implications. To better understand how the BMI-associated NAWM and NAGM brain volume was distributed, we evaluated NAGM and NAWM volumes using the 3 discrete diagnostic classifications, normal BMI, overweight, and obese. The obese group showed a reduction in NAWM brain volume over the 3 years of the study, and the overweight and the normal BMI groups demonstrated fluctuating results in NAWM brain volume measures over the 3 years of the study. The obese and overweight groups showed a reduction in NAGM volume over the 3 years of the study, and the normal BMI group demonstrated a marginal increase in NAGM brain volume over the 3 years of the study (Table 4).

A multivariate linear regression analysis was performed to evaluate the impact of the baseline BMI on the 36th month

### Table 3. Baseline and month 36th summary of changes between baseline and month 36 with effect size (n = 768).

| VARIABLES                  | MEAN DIFFERENCE | POOLED SD | EFFECT SIZE | t VALUE | P VALUE |
|----------------------------|-----------------|-----------|-------------|---------|---------|
| Brain volume in ml         |                 |           |             |         |         |
| NA white matter            | -1.55           | 19.8      | .16         | 1.96    | .031    |
| NA gray matter             | -3.47           | 25.8      | .27         | 3.79    | <.001   |
| Spinal fluid volume        | 9               | 18.3      | .98         | -13.52  | <.001   |
| Cognitive function in points |                |           |             |         |         |
| PASAT (categorical)        | 3.79            | 7.8       | $\chi^2(1) = 37.157$ | .07     | <.001   |
| PASAT (continuous)         |                 |           | .97         | 13.29   | <.001   |
| BMI in kg/m$^2$            |                 |           |             |         |         |
| BMI (continuous)           | .231            | 3.6       | .13         | -1.78   | .075    |
| BMI (categorical)          |                 |           | $\chi^2(1) = 3.41$ | .065    |         |

Note. The brain volume was evaluated based on paired t-tests using SPSS IBM version 23. The BMI and categorical cognitive function (PASAT) were evaluated based on paired t-test (continuous) and McNemar test (categorical). Cohen’s d effect size for 2-tail t-test was calculated based on the absolute value of the mean difference between the baseline and 36th month groups divided by .5 times the pooled standard deviation (Cohen’s d = |m2 − m1|/[(sd1 + sd2)/2], n1 = n2).

### Table 4. BMI and Normal Appearing Brain Volume or Cognitive Function at Baseline and 36th Month (n = 768).

| CHARACTERISTICS | NORMAL BMI | OVERWEIGHT | OBESE     |
|----------------|------------|------------|-----------|
| NAWM Baseline  | 463.67 (54.25) | 475.40 (52.12) | 470.14 (55.92) |
| 36th month     | 462.69 (56.50) | 475.13 (54.18) | 466.19 (62.19) |
| NAGM Baseline  | 581.81 (64.95) | 594.84 (59.70) | 587.21 (64.64) |
| 36th month     | 583.07 (62.12) | 588.74 (56.41) | 583.39 (66.04) |
| Cognitive function Baseline | 50.37 (10.22) | 49.89 (10.23) | 49.82 (10.72) |
| 36th month     | 54.51 (7.53)  | 54.28 (7.68)  | 52.85 (8.51)  |
NAWM and NAGM brain volume. The baseline BMI had no significant effect on the 36th month NAGM brain volume; however, it had a significant effect on the 36th month NAWM brain volume. Therefore, a hierarchical multivariate linear regression model was done to assess the effect of the baseline BMI on NAWM brain volume after accounting for other independent variables noted above. The first block included the independent variable, BMI, the second block included the cardiovascular variables and the third block included MS related factors and demographics (Table 5). The baseline BMI (categorized into normal, overweight, and obese) exhibited an effect on the 36th month NAWM brain volume. Compared to the normal BMI group, the overweight BMI group showed significantly higher NAWM volume (B = 12.3, t = 2.300, P = .022), but the obese group displayed similar white matter volume (B = 4.1, t = .732, P = .432) as the normal BMI group (Table 5). The association between BMI and NAWM was eliminated in the third hierarchical block and sex was shown to be a confounder variable in the relation between BMI and NAWM (Table 5). Each demographic and disease-related variable was tested separately in the Hierarchical model, demonstrating that sex was the confounder between BMI and NAWM. Similarly, ITT analysis of the categorical BMI effect showed that the pooled slope was \(\sim 11\) mL larger volume in NAWM in those with overweight BMI (\(P = .036\)) compared to normal weight, but there was not a difference for the obese group.

Mixed model analysis was performed to evaluate the effect of the baseline BMI as a continuous and as a categorical variable on the longitudinal normal appearing brain volume changes during the 3 years of the study. Thus, the analysis was performed to investigate whether there was an association between the

| TABLE 5. Hierarchical linear regression model for baseline BMI and the 36th month white matter (n = 768). |
|----------------------------------|
| VARIABLES | BASE MODEL | BLOCK 1 | BLOCK 2 – FULL MODEL |
|-----------|------------|---------|----------------------|
|           | F(2, 765) = 2.769, P = .063 | F(6, 761) = 3.264, P = .004 | F(15, 752) = 16.931, P < .001 |
| Constant  | 462.6 | <.001 | 437.742 | <.001 | 495.14 | <.001 |
| BMI       |          |         |         |       |         |         |
| Normal BMI (ref) |          |         |         |       |         |         |
| Overweight | 12.3 | .10 | .022 | 12.8 | .10 | .018 | 3.8 | .030 | .434 |
| Obese     | 4.06 | .03 | .432 | 2.8 | .026 | .597 | 3.4 | .031 | .473 |
| Blood Pressure |          |         |         |       |         |         |
| Systolic BP |       |         |         |       |         |         |
| Diastolic BP |       |         |         |       |         |         |
| Smoking   |          |         |         |       |         |         |
| Ex-smoker | 9.847 | .069 | .096 | 10.338 | .072 | .048 |
| Never smoker | 15.046 | .128 | .002 | 16.25 | .138 | <.001 |
| Current smoker (ref) |          |         |         |       |         |         |
| Disability/EDSS |          |         |         |       |         |         |
| Illness duration |       |         |         |       |         |         |
| Duration from 1st symptom | -1.992 | -.087 | .011 |
| Relapse rate |          |         |         |       |         |         |
| Relapse in last 3yr | -3.285 | -.048 | .137 |
| Baseline age |          |         |         |       |         |         |
| Sex       |          |         |         |       |         |         |
| Female | -59.045 | -.447 | <.001 |
| Male (ref) |          |         |         |       |         |         |
| Race      |          |         |         |       |         |         |
| Race AA | 8.252 | .035 | .480 |
| Race CA | 31.712 | .169 | <.001 |
| Other race (ref) |          |         |         |       |         |         |

Note. Dependent variable = white matter brain volume at 36th month; Duration Dx = duration of illness from diagnosis; SBP = systolic blood pressure; DBP = diastolic blood pressure.
baseline BMI and changes in the outcome, brain volume, over time. The mixed model analysis failed to show an effect of the baseline BMI on NAWM brain volume longitudinally. The mixed model analysis showed a statistically significant effect of BMI as categorical and continuous BMI on the NAGM but showed variant results (Tables 6 and 7). These analyses showed that NAGM brain volume increased .11 mL in patients who had a normal BMI than those who were obese ($F(1, 2683.19) = 5.963, P = .003$) (Table 7) (Figure 1). Similarly, the ITT analysis of the pooled effect of the categorical baseline BMI on NAGM volume showed a NAGM increase of .12 mL ($P = .004$) in patients that had a normal BMI. To evaluate the consistency of these findings with categorical BMI distributions of normal vs high BMI, we also evaluated BMI as a dichotomous predictor (eg, $\geq 25$ vs $< 25$ kg/m$^2$) and found similar results. There was an increase of .13 in NAGM volume in those with normal BMI group compared to the overweight and obese group ($\geq 25$ kg/m$^2$) ($F(1, 683.43) = 11.056, P = .001$). Additionally, the effect of the baseline BMI as a continuous variable demonstrated a NAGM volume decrease of .01 mL with every increase of 1 unit of BMI ($F(1, 2684.13) = 9.435, P = .002$) (Table 6). Comparably, the ITT analysis of the pooled effect of the continuous BMI variable on NAGM brain volume showed a decrease in .01 mL in NAGM volume with every increase of 1 unit of BMI ($P < .001$).

### Discussion

This study found that being overweight predicted an increase in NAWM brain volume as compared with normal BMI, but surprisingly the obese group was not different from the normal BMI group. Thus, overweight patients with RRMS had higher white matter volume (12 mL) than those with normal BMI at 36 months, but it was not a clinically meaningful change. Interestingly, a few non-MS studies have supported the protective effect of high BMI. These studies reported larger regional white matter brain volumes in obese individuals compared to normal weight controls, possibly due to increased density of the lipid-based myelin sheath. Importantly, sex acted as a confounder in the association between BMI and NAWM brain volume.

### Table 6. Mixed model for repeated measures analysis: Baseline BMI/continuous, time, and normal appearing gray matter brain volume longitudinally ($n = 768$).

| PARAMETER                  | $\beta$ | STD. ERROR | DF   | $T$  | $P$ | 95% CONFIDENCE INTERVAL |
|----------------------------|---------|------------|------|------|-----|-------------------------|
|                            |         |            |      |      |     | LOWER BOUND            | UPPER BOUND |
| Intercept                  | 650.54  | 15.64      | 687.06 | 41.602 | 0.000 | 619.84 | 681.24 |
| Baseline BMI               | .41     | .28        | 710.76 | 1.449 | .148 | -.14 | .96   |
| Time                       | .154    | .08        | 2684.28 | 1.879 | .060 | -.007 | .31   |
| Baseline BMI * time        | -.01    | .003       | 2684.74 | -3.035 | .002 | -.01  | -.003 |
| Baseline age               | -1.43   | .22        | 683.07 | -6.580 | <.001 | -1.86 | -1.00 |
| Sex Female Male (ref)      | -69.04  | 4.19       | 678.21 | -16.453 | <.001 | -77.27 | -60.79 |
| Race                       |         |            |      |      |     |            |       |
| African American           | -.524   | 12.07      | 678.67 | -.434 | .664 | -28.94 | 18.46 |
| Caucasian                  | 42.18   | 9.78       | 678.04 | 4.313 | <.001 | 22.97 | 61.38 |
| Others (ref)               |         |            |      |      |     |            |       |
| Smoking                    |         |            |      |      |     |            |       |
| Ex-smoker                  | 8.59    | 5.37       | 678.69 | 1.601 | .110 | -.194 | 19.13 |
| Never smoker               | 15.03   | 4.57       | 678.81 | 3.288 | .001 | 6.05  | 24.00 |
| Current smoker (ref)       |         |            |      |      |     |            |       |
| Time from 1st symptom      | -.37    | .38        | 680.37 | -.968 | .333 | -1.11 | .38   |
| Relapse rate 3yrs          | -.539   | 2.14       | 677.96 | -2.517 | .012 | -9.59 | -1.18 |
| Disability (EDSS)          | -.32    | .38        | 2868.83 | -.830 | .407 | -1.06 | .43   |
| Disease modifying          |         |            |      |      |     |            |       |
| Glatiramer acetate         | -.319   | 4.46       | 678.27 | -.714 | .476 | -11.95 | 5.58  |
| Interferon                 | -2.52   | 4.76       | 678.05 | -.529 | .597 | -11.88 | 6.83  |
| Interferon+Glatiramer (ref)|         |            |      |      |     |            |       |

Dependent Variable: Gray Matter Brain Volume. EDSS = Expanded Disability Status Scale; ref = reference group. The interaction of baseline BMI*Time = $F(1/2684.13) = 9.435, P = .002$. 

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likely attributed to the fact that most of the study participants were females who have different developmental phases.

High BMI was found to have a statistically significant effect on NAGM volume. The continuous and categorical baseline BMI models estimated contradictory effects on the NAGM brain volume changes throughout the 3 years of the study, possibly indicating different cutpoints matter or some lack of fit of the linear model to the data with the continuous BMI. Although these results were statistically significant, their clinical meaningfulness is questionable in view of the lack of consistency, the large sample size and the resulting high power for statistical significance in the analyses. Nonetheless, in prior studies, higher BMI appeared to be associated with similar reductions in gray matter volume and brain parenchymal volume.19-21

This study also found that BMI had no effect on cognitive function as assessed by the PASAT. The majority of the patients in the study were newly diagnosed patients with short duration of illness and high cognitive performance and all were treated with one or more DMTs, which might have been the reason for the absent relationship between BMI and cognitive function. The PASAT scores improved from baseline to month 36th, which could be explained by the data analysis showing that the missing 36th month data were of those with the most impaired cognitive performance who either withdrew from the study or refused to take this difficult test. Similarly to our secondary analysis results, a recent large study (n = 8713) of patients with MS revealed no association between the Processing Speed Test (PST) and BMI as continuous or categorical variable.36 Contrastingly, other cross-sectional evidence has

| PARAMETER | ß    | STD. ERROR | DF | T     | P   | 95% CONFIDENCE INTERVAL |
|-----------|------|------------|----|-------|-----|------------------------|
|           |      |            |    |       |     | LOWER BOUND          | UPPER BOUND |
| Intercept | 665.59 | 14.04    | 679.17       | 47.42  | .000 | 638.04         | 693.15      |
| BMI       |       |            |            |       |     |                       |             |
| Normal BMI| −7.24 | 4.61       | 709.59       | −1.571 | .117 | −16.29        | 1.81        |
| Overweight| −3.42 | 4.72       | 709.35       | −.724  | .470 | −12.60        | 5.86        |
| Obese BMI (ref) time | −.11 | .03        | 2683.57       | −3.627 | <.001 | −.17         | −.05        |
| BMI * time |       |            |            |       |     |                       |             |
| Normal BMI * time | .11 | .05        | 2683.51       | 2.463  | .014 | .02          | .20         |
| Overweight * time | −.04 | .05        | 2683.29       | −.968  | .333 | −.14         | .05         |
| Obese BMI * time (ref) |       |            |            |       |     |                       |             |
| Baseline age | −1.43 | .22        | 682.09       | −6.567 | <.001 | −1.85        | −1.00       |
| Sex       |       |            |            |       |     |                       |             |
| Female    | −68.93 | 4.24      | 677.18       | −16.272 | <.001 | −77.25       | −60.61      |
| Male (ref)|       |            |            |       |     |                       |             |
| Race      |       |            |            |       |     |                       |             |
| African American | −5.22 | 12.07     | 677.64       | −.433  | .665 | −28.92        | 18.48       |
| Caucasian | 42.35 | 9.79       | 677.03       | 4.328  | <.001 | 23.14        | 61.57       |
| Others (ref) |       |            |            |       |     |                       |             |
| Time from 1st symptom | −.38 | .38        | 679.33       | −.994  | .321 | −1.12         | .37         |
| Disability (EDSS) | −.28 | .38        | 2867.21       | −.745  | .456 | −1.03         | .46         |
| Relapse rate in 3yrs | −5.41 | 2.14     | 676.97       | −2.530 | .012 | −9.62        | −1.21       |
| Disease-modifying therapy |       |            |            |       |     |                       |             |
| Glatiramer acetate | −3.29 | 4.47      | 677.26       | −.738  | .461 | −12.07        | 5.48        |
| Interferon | −2.69 | 4.77      | 677.05       | −.564  | .573 | −12.06        | 6.78        |
| Interferon+Glatiramer(ref) |       |            |            |       |     |                       |             |
| Smoking  |       |            |            |       |     |                       |             |
| Ex-smoker | 8.45  | 5.37       | 677.69       | 1.573  | .116 | −2.10        | 19.00       |
| Never smoker | 15.14 | 4.57     | 677.81       | 3.307  | .001 | 6.15         | 24.13       |
| Current smoker (ref) |       |            |            |       |     |                       |             |

Note: The Categorical Baseline BMI * Time = F(2, 2683.19) = 5.963, P = .003. Obese BMI as reference was chosen by the SPSS Mixed model analysis.
found that obesity is linked to reduced cognitive functions, particularly in executive, attention, and memory domains, which are highly prevalent in MS. In addition, although an effect between BMI and cognitive function has been seen mostly in the aging healthy population and partially in a few MS studies, the interaction between BMI and cognitive function in patients with MS, and the role of BMI as a risk factor for cognitive dysfunction are complex and highly debated. However, if an association exists between cognition and BMI, the mechanisms are unclear, and it might be through complex mechanisms that involved depression, exercise, or other factors associated with both obesity and cognitive function.

**Limitations of the Study**

This study had several limitations that warrant discussion. The CombiRx primary study did not include a placebo control arm; therefore the comparisons and findings are related to patients with RRMS on DMTs (glatiramer acetate or interferon beta-1a or combination of both). The lack of age-sex matched control group influenced the ability to ascertain the effects of the disease itself, BMI and/or other covariates on the outcomes. Additionally, the CombiRx study included patients with MS who were newly diagnosed, and therefore had minimal functional (mean EDSS < 2) or cognitive disability. Patients with longer disease duration might have had greater disability that may have affected their BMI, brain volume and cognitive function.

In addition, the follow-up time of this study was 3 years, and while among the longest RRMS trial, still may not have been enough time to detect meaningful changes in BMI, cognitive function, and brain volume that could be found in longer follow-up periods. Furthermore, BMI is often considered to be an inaccurate measure of body fat content and does not take into account muscle mass, bone density, overall body composition, and racial and sex differences. Other measures such as waist circumference, body fat percent, and other reliable methods to measure normal, overweight and obesity conditions may provide a better assessment of their impact on the course of MS. Additionally, as noted above there are some limitations related to the PASAT. Nonetheless, recent evidence supports a correlation of the PASAT with a highly sensitive test like the SDMT that is now often used in MS research and clinical practice. Finally, the exclusion of patients with major comorbidities in the CombiRx study is both a strength and a limitation. While it somewhat limits generalizability to a wider group of individuals with MS, it also helped isolate the effects of obesity separate from other comorbid diseases and their treatments. In addition, other variables, such as physical activity and mood disorders, were not included in the CombiRx trial. These factors have an impact on brain volume and cognitive function in people with MS, which can affect the results of the study.

**Conclusion**

This study showed questionable effects of BMI as a continuous or categorical variable on cognitive function and normal appearing brain volume. Consistently, there was no predictable effect of the BMI on cognitive function as measured by the PASAT. Furthermore, the likelihood of meaningful impact of
BMI on cognition or brain volume in early RRMS patients over 3 years seems remote due to the rigor and sample size of this trial, the consistency between the 3 year completers and the ITT results with imputation. To address some of the issues raised given the mixed nature of the results, future longitudinal prospective research studies should include a few anthropometric measurements and other cognitive tests assessing their associations and the impact of these measurements on the course of MS.

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