Economic burden of multiple sclerosis on Kuwait health care system

Maryam S. Alowayesh1*, Samar F. Ahmed2,3, Jasem Al-Hashel2,4, Raed Alroughani5

1 Department of Pharmacy Practice, School of Pharmacy, Kuwait University, Jabriya, Kuwait, 2 Department of Neurology, Ibn Sina Hospital, Sabah Medical Area, Kuwait, 3 Department of Neurology and Psychiatry, Minia University, Minia, Egypt, 4 Department of Medicine, Faculty of Medicine, Kuwait University, Jabriya, Kuwait, 5 Division of Neurology, Department of Medicine, Amiri Hospital, Sharq, Kuwait

* alowayeshms@hsc.edu.kw

Abstract

Background

Multiple Sclerosis (MS) is a chronic neurological disease with heavy economic and social burdens resulting in significant disability.

Objective

This study aims to (1) measure the cost of health resources utilization by MS patients and (2) to examine the difference in utilization and its attributed costs amongst patients who may have a different course of MS and expanded disability status scale (EDSS) scores.

Methods

A cross-sectional study using Kuwait National MS registry was conducted to estimate the costs of utilization of resources from 2011 to 2015.

Results

Between the period 2011–2015, 1344 MS patients were included in the registry. The average annual cost per MS patient has increased from $10,271 in 2011 to $17,296 in 2015. Utilization of disease-modifying therapies (DMTs) was the main driver of costs reaching 89.9% in 2015. Throughout the five-year period, the occurrence of relapses decreased from 21.8% to 12.2% (p <0.0001). During this same period, ambulatory relapse treatment increased by 5.8% while hospitalizations decreased by 2.6%. Patients with a moderate EDSS score (3.5–6) had the highest average cost (p<0.0001) compared to mild and severe EDSS scores.

Conclusions

Multiple sclerosis has been a significant economic burden on the Kuwait healthcare system. DMTs are the main driver of cost.
Introduction

Multiple Sclerosis (MS) is a chronic debilitating disease with heavy economic and social burdens resulting in severe disability and social dependence.[1] Because of the early onset of MS, it can often occur during the patient’s most productive working years, thus creating potentially large societal costs.[2] In addition to its burden on the patients and society, the entire healthcare system shares the financial burden of MS. [3] As a result of relapses and the progressive nature of MS, patients require repeated hospitalizations during disease exacerbations or worsening of their neurological disabilities. [4] In North-American and European studies, it has been reported that MS patients are more than twice as likely to be hospitalized or to consult a healthcare professional than patients without MS.[5–9]

When compared with the direct medical costs of other chronic conditions described in the literature, MS ranked second behind congestive heart failure. [10] According to several recent MS cost-of-illness studies, direct medical costs accounted for 64–77% of all costs with DMTs being the main driver of cost.[10,11]

In Kuwait, reported prevalence rate of MS was 85.05 per 100,000 persons in 2011.[12] Despite this high prevalence rate, the economic burden associated with the disease in Kuwait, as well as in the Middle East, is unknown. Detailed knowledge of the costs of an illness can provide the essential background that is necessary for policymakers to make informed decisions regarding which areas of disease treatment need to be addressed first and to subsequently set up and prioritize health-care policies and interventions.[13] Economic burden studies have become increasingly important under fast-changing healthcare systems.[3] Therefore, our aim was to measure the cost of health resources’ utilization by MS patients during the period 2011–2015 and to follow up by examining the differences in utilization and attributed costs amongst patients with different EDSS scores.

Methods

Patients and data collection

Ministry of Health (MOH) institutional review board has approved this study. Written consent forms have been obtained from the participants. This cross-sectional study was conducted using data from Kuwait National MS Registry. Established in 2010, this registry accounts for nearly 95% of the MS patients diagnosed in Kuwait.[14] The registry includes the neurology tertiary hospital, other peripheral hospitals that have neurology units, and MS clinics. All patients were assessed by neurologists who are experienced in MS diagnosis using the revised 2010 McDonald diagnostic criteria.[15] Patients were classified either as clinically isolated syndrome (CIS), RR MS, progressive relapsing (PR) MS, secondary progressive (SP) MS, or primary progressive (PP) MS.[16] Patients included in the registry are seen at least twice per year during scheduled visits. A range of laboratory and radiological investigations are routinely ordered depending on the patient’s clinical status and what DMTs have been prescribed. Additionally, unscheduled visits and investigations due to relapses, adverse events, or other medical events are recorded in the registry. The institutional ethical committee approved the study and informed consent forms were obtained from all patients.

Costing

Based on Trisolini and colleagues’ conceptual model for MS costs, this study collected only direct medical costs.[17] Five years of data were collected from 2011–2015. All unit costs were obtained from Ministry of Health personnel. They included inpatient hospital admissions, outpatient visits, laboratory and radiological investigations, and medications. In the Kuwaiti
healthcare system, all Kuwaitis are entitled to free public hospital care and are entitled to free MS-related prescription medications. Therefore, all direct expenditure is paid by the Kuwaiti healthcare system. For those reasons, the costs in this study were measured only for Kuwaitis since non-Kuwaitis are not covered by the Kuwaiti healthcare system. Coverage of the costs for non-Kuwaitis is made through the patient’s helping fund or by the patients themselves. Unit costs of direct medical expenses are summarized in Table 1.

Statistical analysis

Direct medical annual costs were presented from the healthcare perspective. Disability was quantified using the EDSS.[18] Patients were stratified based on the severity of their disability into three groups based on their EDSS score: mild (EDSS 0–3), moderate (EDSS 3.5–5.5), and severe (EDSS 6–9).[19–20] To compare the EDSS groups, a Chi-square test was used for categorical variables (gender, birth country, MS course) and ANOVA (age, age of onset, and duration of disease) or Kruskal-Wallis test (number of relapses and for cost comparisons) for continuous variables. For the duration of disease comparison, a robust version of ANOVA (Welch test) was used as this variable did not fit the homogeneity of variance assumption. For summarizing costs in each year, it was presented as means and confidence intervals (CIs). When calculating CIs, the skewness of the distribution has to be considered. The level of statistical significance was set at $P < .05$.

Pairwise comparison between years performed using Dunn’s (1964) [21] procedure with a Bonferroni correction for multiple comparisons (adjusted p-values are used) was used. Pairwise comparisons z-tests with Bonferroni correction for multiple comparisons were also used to compare the use of different DMTs between the years by type of intake (ex. IV, PO, SC, IM).

To investigate the relationship between total cost and EDSS scores, a set of linear regression models were constructed based on the data of the period 2011–2015. The EDSS independent variable was used as a continuous or an ordinal variable (mild, moderate, and severe groups). A logarithmic transformation of the total healthcare costs data was applied because its distribution is skewed.

A set of mixed-effect models with patients as random intercept and EDSS group (continuous or ordinal variable with three groups), gender, year, age along with age at MS onset, and disease duration as fixed effect were performed. An interaction term between the year and EDSS groups was added to the models. A final mixed-effect model was chosen based on the AICC criterion (Hurvich and Tsai [22]) and multi-collinearity between covariates.

Results

There were 1344 patients included in the study who were recorded in the Kuwait national MS registry during the period 2011–2015 (Table 2). No patients were excluded for incomplete data. The majority were females ($n = 896, 66.7\%$), and Kuwaiti national represented 87.6\% ($n = 1143$) of the studied cohort. The mean age at MS onset was 26.8 ($\pm 8.8$) years with a mean duration of disease of 8.7 ($\pm 6.9$) years. Most patients had RR course ($n = 990, 75.9\%$). Also, the majority of patients had a mild EDSS score (0–3) ($n = 893, 77.2\%$). The relapse rate decreased significantly from 21.8\% in 2011 to 12.2\% in 2015; ($p < 0.0001$).

The total direct medical costs of MS increased significantly ($p < .005$) from year 2011 to 2015 as shown in Fig 1. There was also a significant difference between per-year and per-patient mean values between 2011 and 2012 years ($p = .016$) followed by significant growth with a one-year lag: 2013 being higher than 2011, 2014 higher than both 2011 and 2012, and 2015 being higher than 2013 and all earlier years (all $p < .005$). This can be interpreted as a
Table 1. Unit costs of direct medical expenses.

| Resource                                      | Cost in US dollars*   |
|-----------------------------------------------|-----------------------|
| **Diagnosis at entry and when needed**       |                       |
| CSF                                           | 523.2                 |
| EP                                            | 327.0                 |
| MRI Brain                                     | 621.3                 |
| MRI Whole spine                               | 1046.4                |
| MRI Thoracic                                  | 523.2                 |
| MRI Cervical                                  | 523.2                 |
| **Diagnosis Labs at entry**                   |                       |
| ANA                                           | 41.2                  |
| ENA                                           | 45.5                  |
| Vitamin B12                                   | 32.4                  |
| TSH                                           | 29.4                  |
| Serum ACE                                     | 47.1                  |
| Anti-NMO IgG test (10% of the sample)        | 164.8                 |
| **Diagnosis Labs for DMTs (before start and during the treatment, according to routine schema)** | |
| CBC                                           | 20.6                  |
| RFT                                           | 53.0                  |
| LFT                                           | 47.1                  |
| Urine Microscopy                              | 11.8                  |
| **Treatment (DMTs)**                          |                       |
| Teriflunomide                                 | 1831.2                |
| Interferon beta — 1a (powder)                 | 1308.0                |
| Interferon beta — 1b                          | 1504.2                |
| Fingolimod                                    | 2746.8                |
| Alemtuzumab **                                | 29430, 49050          |
| Interferon beta — 1a (solution)               | 1275.3                |
| Rituximab**                                   | 3924.0                |
| Dimethyl fumarate                             | 2092.8                |
| Natalizumab                                   | 3106.5                |
| Symptomatic treatment (for EDSS>3.5)          |                       |
| Baclofen                                       | 39.2                  |
| Oxybutynin                                     | 52.3                  |
| Fampridine                                     | 719.4                 |
| **Ambulatory treatment**                      |                       |
| Methylprednisolone (3 days course)            | 359.7                 |
| Methylprednisolone (5 days course)            | 555.9                 |
| **Hospital treatment**                        |                       |
| Methylprednisolone (5 days course)            | 392.4                 |
| Plasmapheresis                                | 9810.0                |
| Hospitalization (per day)                     | 712.9                 |
| **Outpatient visits**                         |                       |
| New patients                                  | 130.8                 |

(Continued)
**Table 1.** (Continued)

| Resource            | Cost in US dollars**** |
|---------------------|------------------------|
| Follow-up patients  | 65.4                   |

* price per month
** 5-day course costs $49050 or 3-day course costs $29430, given annually ($4087.5 per month cost for 5-day course; $2452.5 per month cost for 3-day course)
*** 6-months course ($654 per month cost)
**** the exchange rate was used as for 19.01.2017: 1 KD = $3.27 (https://www.xe.com/currencycharts/?from=KWD&to=USD&view=5Y)
*****These costs include personnel costs.

Abbreviations: CSF—Cerebrospinal Fluid, EP—Evoked potential, MRI—Magnetic Resonance Imaging, ANA—antinuclear antibody, ENA—extractable nuclear antigen, TSH—thyroid-stimulating hormone, ACE—angiotensin converting enzyme, NMO—Neuromyelitis optica, CBC—complete blood count, RFT—Renal function test, LFT—Liver Function Test, DMT—Disease Modifying Therapie, EDSS—Expanded Disability Status Scale, MethylPred—Methylprednisolone

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significant trend towards an increase in mean per-patient costs during the 2011–2015 period. Detailed mean per-patient costs have been summarized in **Table 3.**

The main driver of costs were DMTs since their overall share of the total cost has increased significantly from 2011 to 2015 (84.1% to 89.8% respectively; \( p < 0.0001 \)). Excluding hospitalization costs, all other shares of costs remained the same during the years and its (hospitalization) share in total cost decreased significantly from 2011–2015 (3.9% to 0.7% respectively; \( p < 0.0001 \)). Detailed resource utilization for each year was summarized in **Table 4.**

There was a statistically significant (\( p < .05 \)) growth in intravenous (IV) and oral (PO) DMT utilization as shown in **Fig 2.** The share of IV and PO DMTs usage was significantly higher in 2015 compared to 2011 (\( p < .05 \)) and all intermediate shares lie within the 2011 and 2015 values. The shares of intramuscular (IM) and subcutaneous (SC) utilization decreased measurably within the 2011–2015 period (the 2011 and 2015 shares differ significantly (\( p < .05 \)) and all intermediate values lie within these borders).

**Table 5** showed the results of the final regression model selected based on AICC and collinearity criteria (age of onset and interaction between EDSS and years were removed due to colinearity with other covariates). According to these results and after adjustments for confounding factors, patients with mild EDSS level (0–3) had 41% significantly lower costs compared to the severe group. Patients with moderate EDSS level (3.5–5.5) had 31% higher costs compared to the severe group (6–9). The distribution of total costs was not significantly different between males and females (ratio = 1.13; \( p\text{-value} = 0.159 \)). Total costs did increase significantly when disease duration increased (5% increase in total cost for 1-year increase in disease duration) and across years (55%, 34%, 20%, and 17% lower cost for 2011, 2012, 2013, and 2014 compared to 2015). However, total costs decreased noticeably when age increased (2% decrease in total cost for 1-year increase in age).

Analysis of estimated marginal means revealed that the average total cost per patient per year was $16,848 for moderate EDSS group, $12,849 for severe EDSS group, and $7,593 for mild EDSS group (data are not shown in a table).

**Discussion**

Directed towards the healthcare payers, this study provides an insight into the distribution of costs and the resource utilization that is required across direct costs categories and patients
with varying levels of disability. The economic impact of increasing disability (increase in EDSS score) is particularly obvious in the increase in direct costs between mild and moderate disability. Mean annual direct medical costs increased from $7,593 to $16,848 per person with mild and moderate disease, respectively. This increase did not apply to the severe disease category; their annual mean direct medical cost was $12,849. This was in direct contrast to other

| Variables                        | Period       | Total*             | EDSS categories                        | p-value |
|----------------------------------|--------------|--------------------|----------------------------------------|---------|
| Duration of the disease (years)  | 2011–2015    | 8.7 (6.9)          | Mild (0–3)                              | .0005   |
|                                  |              |                    | Moderate (3.5–5.5)                      |         |
|                                  |              |                    | Severe (6–9)                            |         |
| Age of onset (years)             | 2011–2015    | 26.8 (8.8)         | Mild (0–3)                              | .146    |
|                                  |              |                    | Moderate (3.5–5.5)                      |         |
|                                  |              |                    | Severe (6–9)                            |         |
| Female, n (%)                    | 2011–2015    | 865 (66.3)         | Mild (0–3)                              | .001    |
|                                  |              |                    | Moderate (3.5–5.5)                      |         |
|                                  |              |                    | Severe (6–9)                            |         |
| Birth country Kuwait, n (%)      | 2011–2015    | 1114 (87.8)        | Mild (0–3)                              | .670    |
|                                  |              |                    | Moderate (3.5–5.5)                      |         |
|                                  |              |                    | Severe (6–9)                            |         |
| Age (years)                      | 2011–2015    | 33 (10.1)          | Mild (0–3)                              | .0005   |
|                                  |              |                    | Moderate (3.5–5.5)                      |         |
|                                  |              |                    | Severe (6–9)                            |         |
| Duration of the disease (years)  | 2011–2015    | 188 (71.2)         | Mild (0–3)                              | .0005   |
| Age of onset (years)             | 2011–2015    | 44 (16.7)          | Mild (0–3)                              | .0005   |
| Female, n (%)                    | 2011–2015    | 32 (21.1)          | Mild (0–3)                              | .0005   |
| Birth country Kuwait, n (%)      | 2011–2015    | 341 (75.3)         | Mild (0–3)                              | .0005   |
| Age (years)                      | 2011–2015    | 58 (12.8)          | Mild (0–3)                              | .0005   |
| Female, n (%)                    | 2011–2015    | 54 (11.9)          | Mild (0–3)                              | .0005   |
| Birth country Kuwait, n (%)      | 2011–2015    | 436 (80.4)         | Mild (0–3)                              | .0005   |
| Age (years)                      | 2011–2015    | 57 (10.5)          | Mild (0–3)                              | .0005   |
| Female, n (%)                    | 2011–2015    | 49 (9.1)           | Mild (0–3)                              | .0005   |
| Birth country Kuwait, n (%)      | 2011–2015    | 558 (80.6)         | Mild (0–3)                              | .0005   |
| Age (years)                      | 2011–2015    | 60 (8.7)           | Mild (0–3)                              | .0005   |
| Female, n (%)                    | 2011–2015    | 74 (10.7)          | Mild (0–3)                              | .0005   |
| Birth country Kuwait, n (%)      | 2011–2015    | 589 (83.3)         | Mild (0–3)                              | .0005   |
| Age (years)                      | 2011–2015    | 50 (7.1)           | Mild (0–3)                              | .0005   |
| Female, n (%)                    | 2011–2015    | 68 (9.6)           | Mild (0–3)                              | .0005   |
| Birth country Kuwait, n (%)      | 2011–2015    | 135 (10.5)         | MS course, CIS, n (%)                   | .005    |
| Age of onset (years)             | 2011–2015    | 111 (12.7)         | MS course, PP, n (%)                    | .005    |
| Female, n (%)                    | 2011–2015    | 5 (4.3)            | MS course, PR, n (%)                    | .005    |
| Birth country Kuwait, n (%)      | 2011–2015    | 0 (0)              | MS course, RR, n (%)                    | .005    |
| Age (years)                      | 2011–2015    | 121 (9.4)          | MS course, SP, n (%)                    | .005    |
| Female, n (%)                    | 2011–2015    | 121 (9.4)          | Number of relapses                      | .005    |
| Birth country Kuwait, n (%)      | 2011–2015    | 0.13 (.4)          | No relapses, n (%)                      | .005    |
| Age (years)                      | 2011–2015    | 0.15 (.4)          | 1 relapse, n (%)                        | .005    |
| Female, n (%)                    | 2011–2015    | 0.11 (.3)          | 2 relapses, n (%)                       | .005    |
| Birth country Kuwait, n (%)      | 2011–2015    | 0.04 (.2)          |                                        | .005    |
| Age of onset (years)             | 2011–2015    | 0.13 (.4)          |                                        | .005    |
| Female, n (%)                    | 2011–2015    | 0.15 (.4)          |                                        | .005    |
| Birth country Kuwait, n (%)      | 2011–2015    | 0.11 (.3)          |                                        | .005    |
| Age (years)                      | 2011–2015    | 0.04 (.2)          |                                        | .005    |
| Female, n (%)                    | 2011–2015    | 0.13 (.4)          |                                        | .005    |
| Birth country Kuwait, n (%)      | 2011–2015    | 0.15 (.4)          |                                        | .005    |
| Age (years)                      | 2011–2015    | 0.11 (.3)          |                                        | .005    |
| Female, n (%)                    | 2011–2015    | 0.04 (.2)          |                                        | .005    |
| Birth country Kuwait, n (%)      | 2011–2015    | 0.13 (.4)          |                                        | .005    |
| Age (years)                      | 2011–2015    | 0.15 (.4)          |                                        | .005    |
| Female, n (%)                    | 2011–2015    | 0.11 (.3)          |                                        | .005    |
| Birth country Kuwait, n (%)      | 2011–2015    | 0.04 (.2)          |                                        | .005    |

Data are expressed as mean (SD) if not stated.
EDSS: Expanded Disability Status Scale; MS: multiple sclerosis; SD: standard deviation
MS course: CIS: Clinically isolated syndrome, PP: Primary progressive, PR: Progressive relapsing, RR: Relapsing-remitting, SP: Secondary progressive

The shares are calculated from the number of patients with available EDSS information.

p-value for MS course by EDSS group comparison may be invalid as there were many cells with low count

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studies that found that severe disease incurred the highest costs.\cite{11,23} The difference in statistical results can be explained by the fact that the other studies also captured indirect costs like early retirement and productivity loss along with direct non-medical costs associated with disability such as caregiver costs, physiotherapy costs, and transportation. This particular study did not include those additional categories.

The average annual direct medical cost per MS patient has increased from $10,271 in 2011 to $17,296 in 2015. These figures were similar to other studies that looked at direct costs only.\cite{3,10} The average annual cost per MS patient in 2015 was $17,296; this cost is similar to what other countries are spending on MS.\cite{3,10}

A large multi-national cost-of-illness of MS study, done in 16 European countries, was published in 2017.\cite{24} It confirmed that an increase in disability is directly related to an increase in cost.\cite{24} Additionally, it confirmed that cost in severely disabled patients is not driven mainly by direct medical costs. Medical costs only accounted for 26% of the overall cost category. Instead, the rise in costs was primarily a result of the loss of productivity and an accompanying decline in quality of life for those with advancing disability.\cite{24} Productivity decreased from 82% in mildly disabled patients to 8% in severely disabled patients.\cite{24} While

| Table 3. Detailed mean costs per patient by year (in USD). |
|----------------------------------|-------|-------|-------|-------|-------|-------|
|                                 | 2011  | 2012  | 2013  | 2014  | 2015  |
|                                 | Mean  | 95% CI| Mean  | 95% CI| Mean  | 95% CI| Mean  | 95% CI| Mean  | 95% CI|
| Total cost                      | 10271 | (9543–11000) | 12341 | (11553–13130) | 13911 | (13095–14727) | 15424 | (14588–16260) | 17296 | (16464–18127) |
| Diagnosis (CSF + EP + MRI + Diagnosis at entry) | 246 | (215–275) | 338 | (307–374) | 462 | (426–497) | 429 | (399–459) | 379 | (354–405) |
| Diagnosis (DMTs laboratory tests) | 130 | (120–139) | 157 | (147–168) | 180 | (169–191) | 211 | (200–223) | 233 | (222–245) |
| Treatment (DMTs)                | 8639 | (7945–9296) | 10450 | (9791–11217) | 12166 | (11414–12933) | 13442 | (12651–14222) | 15538 | (14773–16357) |
| Symptomatic treatment (EDSS)    | 800 | (635–969) | 1050 | (880–1230) | 903 | (740–1069) | 1059 | (891–1237) | 880 | (743–1031) |
| Hospital treatment              | 401 | (245–572) | 256 | (139–391) | 64 | (18–127) | 118 | (34–243) | 124 | (54–211) |
| Ambulatory treatment            | 17 | (12–24) | 23 | (17–30) | 53 | (44–62) | 66 | (56–76) | 50 | (42–60) |
| Outpatient visits               | 39 | (35–44) | 68 | (63–73) | 83 | (77–89) | 98 | (93–104) | 91 | (86–96) |

CSF: Cerebrospinal fluid; EP–Evoked potential; MRI: Magnetic resonance imaging
DMT: N,N-Dimethyltryptamine; EDSS: Expanded Disability Status Scale

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utility decreased to less than zero in severely disabled patients, fatigue and cognitive difficulties produced a significant impact on utility. A recent study done in the US also found that the percentages of cost increased alongside increased disability. Annual costs per patient were $51,825, $57,889, and $67,116 for mild, moderate, and severe disability, respectively. It is worth noting that the costs of healthcare in the US are significantly higher than other parts of

Table 4. MS resource utilization by year, n (proportion of patients, %).

| Year       | 2011   | 2012   | 2013   | 2014   | 2015   | p-value |
|------------|--------|--------|--------|--------|--------|---------|
| Pairwise comparisons   | A      | B      | C      | D      | E      |         |
| Total Diagnosis        | 290 (31.4) | 378 (36.4) | 540 (47.3) | 642 (52.2) | 640 (49.0) | < .0005 |
| CSF analysis           | 14 (1.5) | 22 (2.1) | 18 (1.6) | 24 (1.9) | 10 (0.8) | 0.069   |
| EP                     | 11 (1.2) | 19 (1.8) | 11 (1.0) | 18 (1.5) | 4 (0.3)  | .007    |
| MRI                    | 243 (26.3) | 344 (33.1) | 497 (43.5) | 617 (50.1) | 630 (48.3) | < .0005 |
| Diagnosis at entry     | 127 (13.7) | 123 (11.8) | 137 (12.0) | 121 (9.8) | 98 (7.5)  | < .0005 |
| Diagnosis (Labs for DMTs) | 467 (50.5) | 566 (54.5) | 642 (56.2) | 723 (58.7) | 828 (63.4) | < .0005 |
| Treatment (DMTs)       | 478 (51.7) | 575 (55.4) | 657 (57.5) | 735 (59.7) | 850 (65.1) | < .0005 |
| Symptomatic treatment (EDSS) | 76 (8.2) | 112 (10.8) | 106 (9.3) | 134 (10.9) | 118 (9.0) | 0.159   |
| Hospital treatment     | 33 (3.6) | 20 (1.9)  | 7 (0.6)   | 8 (0.6)   | 13 (1)   | < .0005 |
| Ambulatory treatment   | 31 (3.4) | 44 (4.2)  | 115 (10.1) | 148 (12.0) | 120 (9.2) | < .0005 |
| Outpatient visits      | 264 (28.5) | 453 (43.6) | 542 (47.5) | 692 (56.2) | 706 (54.1) | < .0005 |

CSF: Cerebrospinal fluid; EP–Evoked potential; MRI: Magnetic resonance imaging; DMT: N,N-Dimethyltryptamine; EDSS: Expanded Disability Status Scale

* Results are based on two-sided tests with significance level 0.05. For each significant pair, the key of the category with the smaller column proportion appears under the category with the larger column proportion

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Fig 2. Utilization of treatments (DMTs) 2011–2015, by type of intake* (%). * IM (intramuscular): Avonex; IV (intravenous): Tysabri, Lemtrada, Rituxan; PO (per os): Aubagio, Tecfidera, Gilenya, SC (subcutaneous): Betaferon, Rebif.

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the world. However, this may be due to the way healthcare is managed and financed within the US.

A recently published systemic review of MS cost-of-illness studies done in OECD (Organization for Economic Co-operation and Development) countries (18 European countries plus the US and Canada) showed that bottom-up costing approach and prevalence approaches were most common, which is the same approach used in this study.[26] Also, it reported that the cost ratios between different severity levels within studies were fairly stable, to the ratio of 1 to 2 to 3 for disability level categories.[26] Further, it mentioned that drugs were the main cost drivers for MS-patients with low disease severity, while the main cost components for groups with more advanced MS symptoms were production losses due to MS and informal care, all of which are similar to the results of our study. [26]

Relapse rate has halved from 2011–2015 (21.8% to 12.2%). This suggests that compliance and possibly efficacy were better with new-generation DMTs.[27–30] Although comparisons across clinical trials is challenging, especially in the absence of head-to-head comparison trials, several retrospective propensity-matched studies showed that new generation DMTs (specifically natalizumab, fingolimod, and alemtuzumab) were associated with a lower risk of relapse compared to the platform therapies (Beta interferon and Glatiramer Acetate) [27–30] Recent data has shown that dimethyl fumarate has a similar efficacy to fingolimod.[31–32] It would be better to state that in the last decade, high efficacy DMTs have emerged with lower relapse rates and MRI activities.

It is also important to note that the establishment of multi-disciplinary MS clinics and the referral to MS specialists resulted in more patients being escalated or switched to high efficacy DMTs and these actions improved the overall adherence rates as per the neurologists’ observation. Both of these factors may have impacted, in a positive way, on the relapse rate in the last few years. In this five-year period, more relapses were treated in ambulatory clinics (85.2% in 2011–92.6% in 2015) and less needed hospitalizations (14.8% in 2011–7.4% in 2015). Treating patients’ relapses in an ambulatory MS clinic costs much less than treating them in a hospital. A relapse that is treated in an ambulatory MS clinic costs around $359–555 per patient, $359 if it was a 3-day course of methylprednisolone, and $555 if it was a 5-day course of

| Parameter          | Ratio* | 95% Confidence Interval | P-value |
|--------------------|--------|-------------------------|---------|
|                    | 95% Lower Bound | 95% Upper Bound |         |
| Intercept          | 21474.08 | 14307.48 | 32230.42 | p < .001 |
| Mild EDSS*         | 0.59 | 0.46 | 0.76 | p < .001 |
| Moderate EDSS*     | 1.31 | 1.01 | 1.71 | 0.043 |
| Maleb              | 1.13 | 0.95 | 1.33 | 0.159 |
| MS Length in years | 1.05 | 1.03 | 1.06 | p < .001 |
| Year = 2011c       | 0.45 | 0.38 | 0.52 | p < .001 |
| Year = 2012        | 0.64 | 0.56 | 0.72 | p < .001 |
| Year = 2013        | 0.80 | 0.72 | 0.90 | p < .001 |
| Year = 2014        | 0.83 | 0.74 | 0.92 | p < .001 |
| Age in years       | 0.98 | 0.97 | 0.99 | p < .001 |

* Ratio = exp (estimate)
* Severe EDSS is reference group;
* Female is reference group; Year = 2015 is reference group

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methylprednisolone. By contrast, an in-hospital treatment of a relapse would cost around $2,371 if it was 3-day admission and $3,919 if it was 5-day admission.

One US study looked at the excess costs that a patient with relapse will have during the year. It compared patients with relapse and patient with no relapse.[33] Patients with relapse were grouped into a low/moderate severity group and the other was the high severity group. [33] The low/moderate severity group and high severity group incurred an excess of $8269 and $24,180 in direct costs compared to the no-relapse group respectively.[33] Another study done in Ireland looked at the direct and indirect cost of an MS relapse.[34] The directs costs of a patient relapse ranged from $469 (low-intensity relapse) to $6,353 (high-intensity relapse). The significant difference between the two sets of costs was mainly caused by hospital admission costs in the high-intensity relapse group, an amount that produced almost 75% of the cost. [34]

From 2011 to 2015, oral DMTs utilization has increased from 5.2% to 27.1% since most of the newer DMTs were oral and more convenient for patient usage. Subsequently, several observational studies showed improvements in adherence with orals compared to injectables. [35–37] Highly efficacious drugs with greater adherence rates provide the greatest real-world effectiveness and may offer the best economic value.[38] However, highly efficacious therapies with low adherence may yield real-world efficacy that is considerably lower than that observed in strictly monitored clinical trials.[38] Moreover, a US study explored the effect of adherence to DMT on the overall spending on MS.[39] It found that adherence to DMT notably reduced the possibility of relapse by 42%, hospitalization by 52%, and emergency visits by 38% (all, \( P < 0.0001 \)). [39] Adherent patients would be predicted to have on average 0.7 fewer outpatient visits each year versus non-adherent patients (\( P < 0.0001 \)). Based on the differences in predicted mean costs, adherence (vs non-adherence) would decrease the total annual medical care costs by $5,816 per patient, including hospitalization costs by $1,953, emergency visits by $171, and outpatient visits by $2,802. [39]

**Limitations**

This study only examined the direct medical costs of MS. Looking at both direct and indirect costs of MS would give a more comprehensive picture of the burden of the disease. A future study is planned to survey patients and explore indirect costs, the patients’ quality of life, and the patients’ adherence to newer DMTs. Moreover, for this particular study, some unit costs of laboratory procedures were hard to obtain since the Ministry of Health (MOH) only had the cost of the machine and reagent. To overcome that limitation, laboratory personnel were consulted to give an estimate of some of these laboratory procedures. In addition, micro-costing an MS hospitalization was challenging. As a result, the approach that was used was to get the estimate from MOH for a neurology-related hospital admission and then add the cost of specific MS drugs and diagnostics.

This study is a cross-sectional study, which means that it is estimating the cost at this point in time. This time-related factor may bias some of the results since patients may be switched from one DMT to another within a short period due to disease reactivity. Additionally, given the referral bias to MS clinics, our findings cannot be generalized as the main source of cost was driven by the DMT prescriptions that are dependent on the treatment protocol used in our center.

**Conclusion**

Multiple sclerosis continues to be a significant economic burden on the Kuwait healthcare system. Disease-modifying therapies seem to be the main driver of cost. Over recent years, oral
and infusion therapies (new-generation DMTs) have been prescribed more often and, as a result, the overall relapse rates have decreased.

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**Author Contributions**

- **Conceptualization:** Maryam S. Alowayesh.
- **Data curation:** Maryam S. Alowayesh.
- **Formal analysis:** Maryam S. Alowayesh.
- **Investigation:** Maryam S. Alowayesh, Samar F. Ahmed, Jasem Al-Hashel, Raed Alroughani.
- **Methodology:** Maryam S. Alowayesh.
- **Project administration:** Maryam S. Alowayesh.
- **Resources:** Samar F. Ahmed, Jasem Al-Hashel, Raed Alroughani.
- **Supervision:** Maryam S. Alowayesh.
- **Visualization:** Maryam S. Alowayesh.
- **Writing – original draft:** Maryam S. Alowayesh.
- **Writing – review & editing:** Maryam S. Alowayesh, Raed Alroughani.

**References**

1. Patwardhan MB, Matchar DB, Samsa GP, McCrory DC, Williams RG, Li TT. Cost of multiple sclerosis by level of disability: a review of literature. Mult Scler 2005; 11:232–239. [https://doi.org/10.1191/1352458505ms1137oa](https://doi.org/10.1191/1352458505ms1137oa) PMID: 15794399
2. Simmons RD. Life issues in multiple sclerosis. Nat Rev Neurol 2010; 6:603–610. [https://doi.org/10.1038/nrneuro.2010.143](https://doi.org/10.1038/nrneuro.2010.143) PMID: 20856267
3. Naci H, Fleurence R, Birt J, Duhig A. Economic Burden of Multiple Sclerosis: A Systematic Review of the Literature. Pharmacoeconomics 2010; 28(5):363–379. [https://doi.org/10.2165/11532230-000000000-00000](https://doi.org/10.2165/11532230-000000000-00000) PMID: 20402540
4. Kobelt G, Pugliatti M. Cost of multiple sclerosis in Europe. Eur J Neurol 2005; 12(Supp. 1):S63–S67.
5. Pohar SL, Jones CA, Warren S, Turpin KV, Warren K. Health status and health care utilization of multiple sclerosis in Canada. Can J Neurol Sci 2007; 34(2):167–174. PMID: 17598593
6. Pope GC, Urato CJ, Kulas ED, Kronick R, Gilmer T. Prevalence, expenditures, utilization, and payment for persons with MS in insured populations (provisional record). Neurology 2002; 58:37–43. PMID: 11781403
7. Gottberg K, Einarsson U, Fredrikson S, von Koch L, Holmqvist LW. Multiple sclerosis in Stockholm County: a pilot study of utilization of health-care resources, patient satisfaction with care and impact on family caregivers. Acta Neurol Scand 2002; 106(5):241–247. PMID: 12371915
8. Beckerman H, van Zee IE, de Groot V, van den Bos GA, Lankhorst GJ, Dekker J. Utilization of health care by patients with multiple sclerosis is based on professional and patient-defined health needs. Mult Scler 2008; 00:1–11.
9. Stolp-Smith KA, Atkinson EJ, Campion ME, O’Brien PC, Rodriguez M. Health care utilization in multiple sclerosis: a population-based study in Olmsted County, MN. Neurology 1998; 50(6):1594–1600. PMID: 9639699
10. Adelman G1, Rane SG, Villa KF. The cost burden of multiple sclerosis in the United States: a systematic review of the literature. J Med Econ 2013; 16(5):639–647. https://doi.org/10.3111/13696998.2013.778268 PMID: 23425293

11. Kanavos P, Tinelli M, Efthymiadou O, Visintin E, Grimaccia F, Mossman J. Towards better outcomes in multiple sclerosis by addressing policy change: The International MultiplE Sclerosis Study (IMPrESS). London School of Economics. UK. 2016

12. Alroughani R, Ahmed SF, Bebabanai R, Khan R, Thussu A, Alexander KJ. Increasing prevalence and incidence rates of multiple sclerosis in Kuwait. Mult Scler 2014; 20(5):543–547. https://doi.org/10.1177/1352458113504328 PMID: 24025709

13. Changik Jo. Cost-of-illness studies: concepts, scopes, and methods. Clinical and Molecular Hepatology 2014; 20:327–337. https://doi.org/10.3350/cmh.2014.20.4.327 PMID: 25548737

14. Alroughani R1, Ashkanani A, Lamdhad S. Clinical characteristics of multiple sclerosis in Kuwait: data from the new MS registry of Amiri Hospital. Int J Neurosci 2012 Feb; 122(2):82–7. https://doi.org/10.3109/00207454.2011.630543 PMID: 21985585

15. Polman CH, Reingold SC, Banwell B, Clanet M, Cohen JA, Filippi M, et al. Diagnostic criteria for multiple sclerosis: 2010 revisions to the McDonald criteria. Ann Neurol 2011; 69:292–302. https://doi.org/10.1002/ana.22366 PMID: 21387374

16. Lublin FD and Reingold SC. Defining the clinical course of multiple sclerosis: results of an international survey. National Multiple Sclerosis Society (USA) Advisory Committee on Clinical Trials of New Agents in Multiple Sclerosis. Neurology 1996; 46:907–911. PMID: 8780061

17. Kurtzke JF. Rating neurologic impairment in multiple sclerosis: an Expanded Disability Status Scale (EDSS). Neurology 1983; 33(11):1444–1452. PMID: 6685237

18. Trisolini M, Honeycutt A, Wiener J, Lesesne S. Global Economic Impact of Multiple Sclerosis: A Literature Review. Multiple Sclerosis International Federation (UK) and RTI International (USA). 2010.

19. Kalincik T, Cutter G, Spelman T, Jokubaitis V, Havrdova E, Horakova D, et al. Brain 2015; 138(Pt 11):3287–98. https://doi.org/10.1093/brain/awv258 PMID: 26359291

20. Kister I, Chamot E, Salter AR, Cutter GR, Bacon TE, Herbert J. Disability in multiple sclerosis: a reference for patients and clinicians. Neurology 2013; 80:1018–1024. https://doi.org/10.1212/WNL.0b013e3182872855 PMID: 23427319

21. Dunn OJ. Multiple comparisons using rank sums. Technometrics 1963; 6:241–252.

22. Hurvich CM, Tsai CL. Bias of the corrected AIC criterion for under fitted regression and time series models. Biometrika 1991; 78:3:499–509.

23. Torabipour A, Asl ZA, Majdinasab N, Ghasemzadeh R, Tabesh H, Arab M. A Study on the Direct and Indirect Costs of Multiple Sclerosis Based on Expanded Disability Status Scale Score in Khuzestan, Iran. International Journal of Preventive Medicine 2014; 5(8):1131–1138. PMID: 25317296

24. Ernstsson O, Gyllensten H, Alexanderson K, Tinghög P, Friberg E, Norlund A. Cost of Illness of Multiple Sclerosis—A Systematic Review. PLoS ONE 2016; 11(7):e0159129. https://doi.org/10.1371/journal.pone.0159129 PMID: 27411042

25. Kalincik T, Brown JWJ, Robertson N, Willis M, Scolding N, Rice CM, et al. Treatment effectiveness of alemtuzumab compared with natalizumab, fingolimod, and interferon beta in relapsing-remitting multiple sclerosis: a cohort study. Lancet Neurol 2017; 16(4):271–281. https://doi.org/10.1016/S1474-4422(17)30007-8 PMID: 28209331

26. Spelman T, Kalincik T, Jokubaitis V, Zhang A, Pellegrini F, Wiendl H, et al. Comparative efficacy of first-line natalizumab vs IFN-β or glatiramer acetate in relapsing MS. Neurol Clin Pract 2016; 6(2):102–115. https://doi.org/10.1212/CPJ.0000000000000227 PMID: 27104064

27. Braune S, Lang M, Bergmann A; NeuroTransData Study Group. Efficacy of fingolimod is superior to injectable disease-modifying therapies in second-line therapy of relapsing-remitting multiple sclerosis. J Neurol 2016; 263(2):327–333. https://doi.org/10.1007/s00415-015-7970-6 PMID: 26645389

28. He A, Spelman T, Jokubaitis V, Havrdova E, Horakova D, Trojano M, et al. Comparison of switch to fingolimod or interferon beta/glatiramer acetate in active multiple sclerosis. JAMA Neurol 2015; 72(4):405–413. https://doi.org/10.1001/jamaneurol.2014.4147 PMID: 25665031
31. Fox RJ, Chan A, Zhang A, Xiao J, Levison D, Lewin JB, et al. Comparative effectiveness using a matching-adjusted indirect comparison between delayed-release dimethyl fumarate and fingolimod for the treatment of multiple sclerosis. Curr Med Res Opin 2017; 33(2):175–183. https://doi.org/10.1080/03007995.2016.1248380 PMID: 27733070

32. Wicks P, Rasouliyan L, Katic B, Nafees B, Flood E, Sasané R. The real-world patient experience of fingolimod and dimethyl fumarate for multiple sclerosis. BMC Res Notes 2016; 9(1):434. https://doi.org/10.1186/s13104-016-2243-8 PMID: 27604188

33. Parisé H, Laliberté F, Lefebvre P, Duh MS, Kim E, Agashivala N, et al. Direct and indirect cost burden associated with multiple sclerosis relapses: Excess costs of persons with MS and their spouse caregivers. J Neurol Sci 2013; 330:71–77. https://doi.org/10.1016/j.jns.2013.04.007 PMID: 23647840

34. O’Connell K, Kelly SB, Fogarty E, Duggan M, Buckley L, Hutchinson M, et al. Economic costs associated with an MS relapse. Multiple Sclerosis and Related Disorders 2014; 3:678–683. https://doi.org/10.1016/j.msard.2014.09.002 PMID: 25891546

35. Bayas A, Mäurer M. Teriflunomide for the treatment of relapsing-remitting multiple sclerosis: patient preference and adherence. Patient Prefer Adherence 2015; 9:265–274. https://doi.org/10.2147/PPA.S61651 PMID: 25709412

36. Bergvall N, Petrilla AA, Karkare SU, Lahoz R, Agashivala N, Pradhan A, et al. Persistence with and adherence to fingolimod compared with other disease-modifying therapies for the treatment of multiple sclerosis: a retrospective US claims database analysis. J Med Econ 2014; 17(10):696–707. https://doi.org/10.3111/13696998.2014.940422 PMID: 25019581

37. Agashivala N, Wu N, Abouzaid S, Wu Y, Kim E, Boulanger L, et al. Compliance to fingolimod and other disease-modifying treatments in multiple sclerosis patients, a retrospective cohort study. BMC Neurol 2013; 13:138. https://doi.org/10.1186/1471-2377-13-138 PMID: 24093542

38. Brandes D, Raimundo K, Agashivala N, Kim E. Implications of Real-world Adherence on Cost-effectiveness Analysis in Multiple Sclerosis. J Med Econ 2013; 16(4):547–551. https://doi.org/10.3111/13696998.2013.774281 PMID: 23391123

39. Burks J, Marshall TS, Ye X. Adherence to disease-modifying therapies and its impact on relapse, health resource utilization, and costs among patients with multiple sclerosis. ClinicoEconomics and Outcomes Research 2017; 9:251–252. https://doi.org/10.2147/CEOR.S130394 PMID: 28496344