Characterizing the Occurrence of Key Clinical Milestones in Duchenne Muscular Dystrophy in the United States Using Real-World Data

Shelagh M. Szabo, Alexa C. Klimchak, Christina Qian, Susan Iannaccone, Evan Popoff and Katherine L. Gooch

Broadstreet Health Economics & Outcomes Research, Vancouver BC, Canada
Sarepta Therapeutics, Inc, Cambridge MA, USA
University of Texas Southwestern, Harry Hines Blvd, Dallas TX, USA

INTRODUCTION

Duchenne muscular dystrophy (DMD) is a severe X-linked progressive neuromuscular degeneration caused by mutations in the gene for dystrophin, a protein required for the structural integrity of muscle cells [1–4]. Affected patients typically present in early childhood with gait abnormalities, muscle weakness, and delayed motor and cognitive function [1, 5–8]. Several well-described clinical cohorts in the United States (US) have documented the inexorable progression of DMD, including from the Cooperative International Neuromuscular Research Group (CINRG), [9] Duchenne Registry, [10] and
Centers for Disease Control and Prevention’s Muscular Dystrophy Surveillance, Tracking, and Research Network (MD STARnet) [11]. Although there is heterogeneity in the exact timing of events, patients inevitably experience key clinical milestones in their progression. Muscular weakness leads to loss of ambulation (LOA) and scoliosis in late childhood [12–14]. Loss of strength in active breathing muscles contributes to respiratory insufficiency and the need for ventilation in the teenage years [15]. Cardiomyopathy also develops in late adolescence and the progression of DMD culminates with early mortality in the third or fourth decade of life [13, 16–19].

While estimates of the frequency and approximate timing of key clinical milestones are available from these well-defined US clinical cohorts, [9–11] estimates from generalizable population-based real-world cohorts to characterize the progression experienced by those with DMD in routine clinical practice are limited. Administrative claims studies are based on healthcare usage data that document patient diagnoses, procedures performed, medications dispensed, physicians visited, and inpatient stays. Such data are often used to characterize patient populations, understand treatment patterns, and assess aspects of the clinical or economic burden of a health condition from a large population perspective [20]. Claims database assessments for DMD however are limited. At the time of this study, the only published claims-based study providing estimates of the clinical burden of DMD was prior to the widespread use of corticosteroids, and included a small number of patients from one commercial plan only [21]. The objective of this study was to estimate the prevalence of key clinical milestones by age among commercially-insured patients with DMD in the US using real-world data.

**MATERIALS AND METHODS**

This study was a US-based, real-world, retrospective cohort study to describe the clinical course of patients with DMD, treated under commercial health insurance plans. The study cohort and entry criteria were described previously in a study examining the characteristics of and economic burden among those with DMD in the US [22].

**Data source**

Data were derived from the IBM MarketScan commercial databases, [23] a set of large, nationally representative healthcare databases. The commercial database contains data for employer-sponsored, privately insured employees and their families, [23] and a total of 78,371,462 unique individuals were covered within the 5-year period from 2013 to 2018. These data have been widely validated for clinical, pharmacoepidemiological and pharmacoeconomic research [24–32].

**Study sample**

MarketScan claims (2013–2018) were used to identify the eligible DMD population, which included all males ≤30 years of age, with a muscular dystrophy (MD) International Classification of Diseases (ICD)-9 code 359.1, MD-related ICD-10 code G71.0, or the Becker/Duchenne MD-specific ICD-10 code G71.01, in any position on ≥2 outpatient DMD medical claims (with ≥30 days between claims) or as the primary or secondary diagnosis of ≥1 inpatient claim; or, a dispensation for eteplirsen. To remove those likely to have other congenital dystrophies, individuals with the following were excluded (see Table A1 and A.2 for applicable codes): [21] ≥2 medical claims for ventilator use separated by at least 180 days before age 6 years; ≥1 medical claim with a Current Procedural Terminology (CPT) code for an orthopedic procedure on the foot before age 3 years; ≥1 medical claim for a power, power-assist, and/or manual wheelchair before age 5 years; ≥1 medication fill (NDC 64406005801) or an injection code (HCPCS J2326) for nusinersen at any point during the study period.

To understand the impact of both follow-up time and age on the observation of relevant outcomes, patients were stratified by age at cohort entry (8 to 10 years, 11 to 13 years, 14 to 16 years, and 17 to 19 years). Within these stratified analyses, to ensure some amount of follow-up was available on all of the patients contributing to these analyses, a one-year minimum follow-up requirement was imposed.

**Study design**

The identification period for study enrollment was April 1, 2013 to March 31, 2018. All DMD cohort members were enrolled at their index visit, the first eligible inpatient or outpatient visit with a relevant diagnostic code or dispensation for the DMD-specific medication, eteplirsen. All cohort members were followed until death (if known), deregistration, or the
end of the follow-up period unless otherwise specified in analyses.

Identifying key clinical milestones

Key clinical milestones of interest included LOA, scoliosis, cardiomyopathy, respiratory involvement, and neurologic or neuropsychiatric involvement. As there is no widely-used or well-validated diagnosis code for LOA, it was identified via the proxy outcome of procedural codes for wheelchair use. Scoliosis was identified by either diagnosis codes for scoliosis or procedural codes for spinal surgery. Cardiomyopathy was identified by diagnosis codes for cardiomyopathy or heart failure, or any dispensions for angiotensin-converting enzyme (ACE) inhibitors, angiotensin II receptor blockers (ARBs), beta-blockers, and/or diuretics (spironolactone or eplerenone). For respiratory involvement, diagnostic codes for respiratory failure as well as procedural codes for tracheostomy, assisted ventilation, and selected codes for pulmonary management were used; more severe respiratory conditions were identified by the subset of diagnostic codes for respiratory failure and the procedural code for tracheostomy only. Finally, identification of neurologic and/or neuropsychiatric involvement was based on diagnosis codes for attention deficit disorders, learning disabilities, pervasive developmental disorders, or behavioral disorders; and procedural codes for neuropsychological testing. See Appendix Table A.2 for details of how the clinical outcomes were identified.

Analysis

To describe the study sample, consistent with the previous analyses, [22] the demographic characteristics were summarized, and median duration of follow-up estimated. Health status over the follow-up period was summarized by the percentage observed with at least one prescription for corticosteroids; comorbidity burden (using the median with interquartile range [IQR] Elixhauser Index score, [33] and the frequency of individual Elixhauser comorbidities); and the prevalence of other key comorbidities of interest. These key comorbidities were identified based on literature review, and including respiratory infectious diseases, anxiety, asthma, depression, fractures/osteoporosis, epilepsy, diabetes mellitus and cystic fibrosis. See Appendix Table A.2 for codes.

Primary analyses

In the primary (non-age-restricted) analyses, the percentages of patients who experience key clinical milestones were tallied and the median (IQR) ages at the first observation during the study timeframe of each outcome were calculated within the available follow-up. Due to the limited follow-up available, incidence of events could not be ascertained as the window of observation per individual will impact whether or not an initial event is observed (Appendix Figure A.1).

Age-restricted analyses

To understand the impact of both follow-up time and age on the likelihood of capturing prevalence of patients experiencing key clinical milestones in each cohort, age-restricted analyses were performed. From the cohorts identified by those restrictions, the percentage of patients who experienced LOA, scoliosis, and neurologic and/or neuropsychiatric involvement was estimated among those aged 8 to 10 and 11 to 13 at cohort entry; and the percentage experiencing cardiomyopathy, respiratory involvement, or severe respiratory outcomes among those 14 to 16 and 17 to 19 years at cohort entry. These age categories were selected for clinical relevance, based on evidence from published studies on the mean age at occurrence of LOA, [34, 35] respiratory involvement, [36, 37] and cardiomyopathy in the US. [13, 36, 37] Estimates of the prevalence of key clinical milestones was compared between the age-restricted cohorts and the overall cohort. Age-restricted Kaplan-Meier (KM) analyses were also conducted to account for censoring to better understand the prevalence of events captured by follow-up time available.

Sensitivity analyses

The impact of key design assumptions on the results were tested in sensitivity analyses. Two age-restricted sensitivity analyses were performed. Firstly, the minimum continuous follow-up requirement of one year in the primary age-restricted analysis was removed, to minimize impact of any time-related biases (i.e requiring a minimum of one-year of follow-up could help contribute to higher observed rates of clinical outcomes). Secondly, the observation window was reduced to just one year, without a minimum continuous follow-up requirement. This was to allow for an estimation of the lower-end of the potential range of estimates of the
prevalence of key clinical milestones among age-restricted patients, to better understand the impact of follow-up time.

To increase the likelihood of eliminating uncertain DMD cases, the criteria for identifying DMD was tested by applying three definitions: 1) restricting the cohort to those <18 years old at index, to eliminate individuals with other potential MDs 2) adding an inclusion criterion of having ≥1 diagnosis from a specialist, and 3) adding an exclusion criterion of having ≥2 claims for any of the following: myoneural disorder (358.x or G70.x), Guillain-Barre syndrome (357.0 or G61.0), or hereditary motor and sensory neuropathy (356.2 or G60.0). For all of these sensitivity analyses around DMD definitions, the median age at and proportion experiencing key clinical milestones was tabulated and compared to those of the primary analyses.

Data in the MarketScan commercial databases are de-identified and are compliant with the Health Insurance Portability and Accountability Act (HIPAA) regulations, thus, Institutional Review Board approval was not required to conduct this study.

RESULTS

Patient characteristics and health status

Table 1 summarizes patient characteristics and health status over the study period. In total, 1,964 patients with DMD were identified in the primary analysis. Median (IQR) age at cohort entry was 15 (9–21) years, with a median of 1.7 years of follow-up. At least one corticosteroid dispensation was observed among 38.8% over available follow-up; additional breakdown of corticosteroid use by age group at cohort entry is included in the appendix (Appendix Figure A.2). Over the study period, the median (IQR) unweighted Elixhauser score was 1 (0–3), and cardiac and pulmonary comorbidities were the most frequent individual Elixhauser comorbidities observed. Of the other key comorbidities, respiratory infectious disease was the most common (observed in 48.8% of the cohort) followed by anxiety disorders (15.4%) and asthma (14.5%).

The prevalence of key clinical milestones

Among the DMD cohort from the primary analysis, neurologic and/or neuropsychiatric involvement was observed among 27%, LOA among 45%, scoliosis among 30%, cardiomyopathy among 46%, and respiratory involvement among 34%. The ages at first observed claim for each key clinical milestone in this cohort are presented in Fig. 1. The median (IQR) age at neurologic and/or neuropsychiatric involvement was first recorded at 11 (8–16) years old; LOA was 16 (12–22) years old; for scoliosis, 16 (12–20) years old; for cardiomyopathy, 17 (13–22) years old; and respiratory involvement, 19 (15–24) years old. Age-restricted analyses were also performed to estimate the occurrence of those key outcomes stratified by age at cohort entry. Compared to the overall cohort, the prevalence of key clinical milestones was higher among the age- and time-restricted subset (Table 2), as expected given that the age and time restrictions aimed to focus the observation window on the time period where these outcomes were more likely to have first occurred. As the higher observed prevalence could also be due, in part, to the minimum follow-up requirement, a variation of this analysis was explored where the minimum follow-up requirement was removed (sensitivity analysis #1), and varied to explore the impact of a shorter observation window (sensitivity analysis #2). Due to the nature of age-restriction, the median age at outcome was largely correlated with the respective age restriction, across all scenarios.

The age-restricted KM analyses showed that as follow-up (time from cohort entry) increases, the prevalence of events captured increases (Table 3). By year one, approximately 25% of patients aged 8 to 10 years at cohort entry had an event of LOA captured, compared to 40% by year two, and over 50% by year three. The corresponding values were higher among those aged 11 to 13 years, ranging from approximately 40% by year one to 60% by year three. Similar trends are observed for other key clinical milestones, with a few showing trends of plateau between year two and three, such as respiratory involvement among those aged 17 to 19 years at index which started to plateau at approximately 42% by the end of year two.

Additional sensitivity analyses

In the sensitivity analysis where the criterion for a minimum of one-year of continuous follow-up was removed, the observed percentage experiencing key clinical milestones was less than in the age-restricted analyses. However, estimates generally remained higher than in the primary analyses (Table 2). In addition, in the sensitivity analysis where the observation window was shortened to one year,
the observed percentage experiencing key clinical milestones was expectedly even lower than other scenarios explored.

The sensitivity analyses exploring different DMD cohort definitions (i.e. restricting to patients < 18 years old at index, requiring patients to have at least one diagnosis from a specialist, or excluding patients with myoneural disorder, Guillain-Barre syndrome, or hereditary motor and sensory neuropathy) showed that patients identified by these definitions had similar durations of follow-up, and frequency of corticosteroid use compared to the primary analyses cohort. Age at index, by definition, was lower among the cohort < 18 years old at index, however remained comparable across the other sensitivity analysis cohorts. The frequency and age at key clinical milestones were also comparable between the sensitivity and primary analyses, with the exception of the cohort who was < 18 years old at index, which by definition would have lower estimated median ages at the occurrence of these comorbidities due to the elimination of older DMD patients. Among the cohort < 18 years old at index, the overall frequency of patients with a record of LOA was comparable to
that of the base case definition; median ages were higher for outcomes occurring earlier in patients’ lives, and lower for outcomes that tend to occur in the later stages (Appendix Table A.3).

**DISCUSSION**

Large well-conducted clinical studies and registries have documented the characteristics of the clinical progression of DMD; [12, 13, 18, 19, 39] but how these align with estimates based on real-world data has not previously been reported. This health insurance claims study used the large representative MarketScan commercial database to estimate the prevalence of notable clinical milestones that characterize the natural history of DMD, by age. Approximately 2,000 patients with commercial insurance coverage in the US were included.

The results of the primary (non-age-restricted) analyses showed a slightly older median age at key clinical milestones compared with published estimates from clinical studies, [12, 13, 18, 19, 39] which suggested that at least for some patients, the events captured were potentially not the initial diagnosis related to that milestone. However, as is expected given the relatively short follow-up available in the dataset, the observed frequency of patients experiencing each milestone was less than anticipated overall. As a result, age-restricted analyses were performed. These analyses generally showed higher frequencies of key clinical milestones compared to the primary analysis, as expected given that the age and time restrictions aimed to focus the observation window on the time period where these outcomes were more likely to have occurred. The frequency of LOA among children and young teenagers in the present study – observed in 55% – was consistent with published estimates of the age at LOA range from 30% by 10 years [12] through 95% by 15 years of age [13]. The frequency of scoliosis, documented in up to 50% of young teenagers over their first two years of follow up, was also consistent with published estimates suggesting 60% of DMD patients have scoliosis by 15 years of age [13]. Neurologic and/or neuropsychiatric involvement was observed in less than half of children and young teenagers in the current study, which is also broadly consistent with published estimates of the age at diagnosis of common neuropsychiatric complications in DMD; [40] unlike LOA, for example, it is not necessarily to be expected that these complications would be identified in everyone with DMD [41]. Respiratory involvement was observed in almost half of the cohort of older teenagers, and published estimates suggest 40% to 50% of patients require ventilation over 20 years of age [39, 42]. Finally, estimates of cardiomyopathy and heart failure were on the lower end of published estimates of 68% to 93% diagnosed with cardiomyopathy by 20 years of age [13, 18, 19].

The potential challenges in ascertaining clinical outcomes when using administrative data warrant further discussion. The MarketScan databases are based on claims submitted for reimbursement and not research purposes, and billing practices vary between providers and by insurance type. In the current study, identification of scoliosis, neurologic involvement, cardiomyopathy, respiratory failure, and heart failure primarily relied on diagnosis codes. However, the presence of a diagnostic code alone cannot indicate...
**Table 2**

The percentage with a record of a key clinical milestone, and median age per milestone, among patients with DMD stratified by age, follow-up, and observation window

<table>
<thead>
<tr>
<th>Age at cohort entry</th>
<th>Median (IQR) follow-up</th>
<th>Key clinical milestone</th>
<th>n(%)*</th>
<th>Median (IQR) age at outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>LOA</td>
<td>112 (55.4)</td>
<td>10 (10 to 11)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Scoliosis</td>
<td>76 (37.6)</td>
<td>10 (10 to 11)</td>
</tr>
<tr>
<td>8 to 10 years (n = 202, 10.3% of total cohort)</td>
<td>2.0 (1.7 to 2.0)</td>
<td>Neurologic/neuropsychiatric involvement</td>
<td>86 (42.6)</td>
<td>10 (9 to 10.75)</td>
</tr>
<tr>
<td>11 to 13 years (n = 188, 9.6% of total cohort)</td>
<td>2.0 (1.7 to 2.0)</td>
<td>LOA</td>
<td>104 (55.3)</td>
<td>13 (12 to 14)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Scoliosis</td>
<td>97 (51.6)</td>
<td>13 (12 to 14)</td>
</tr>
<tr>
<td>14 to 16 years (n = 198, 10.1% of total cohort)</td>
<td>2.0 (1.8 to 2.0)</td>
<td>Neurologic/neuropsychiatric involvement</td>
<td>77 (41.0)</td>
<td>13 (12 to 14)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Respiratory involvement</td>
<td>89 (44.9)</td>
<td>16 (13 to 16)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Severe respiratory outcomes</td>
<td>47 (23.7)</td>
<td>16 (15 to 17)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Cardiomyopathy</td>
<td>135 (68.2)</td>
<td>16 (15 to 16)</td>
</tr>
<tr>
<td>17 to 19 years (n = 129, 6.7% of total cohort)</td>
<td>2.0 (1.6 to 2.0)</td>
<td>Respiratory involvement</td>
<td>59 (45.7)</td>
<td>18 (17 to 19)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Severe respiratory outcomes</td>
<td>29 (22.5)</td>
<td>18 (17 to 20)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Cardiomyopathy</td>
<td>75 (58.1)</td>
<td>18 (17 to 18)</td>
</tr>
</tbody>
</table>

Sensitivity analysis variation #1: Two-year observation window; no minimum follow-up imposed

|                      |                        | LOA                            | 109 (42.1) | 10 (9 to 11)  |
|                      |                        | Scoliosis                      | 65 (25.1)  | 10 (9 to 11)  |
|                      |                        | Neurologic/neuropsychiatric involvement | 95 (36.7)  | 9 (9 to 10)  |
| 8 to 10 years (n = 258, 13.2% of the total cohort) | 2.0 (1.1 to 2.0) | LOA                            | 114 (47.9) | 13 (12 to 13) |
| 11 to 13 years (n = 238, 12.1% of the total cohort) | 2.0 (1.2 to 2.0) | Scoliosis                      | 97 (40.8)  | 13 (12 to 13) |
| 14 to 16 years (n = 281, 14.3% of the total cohort) | 2.0 (0.9 to 2.0) | Neurologic/neuropsychiatric involvement | 79 (33.2)  | 13 (12 to 13) |
| 17 to 19 years (n = 174, 8.8% of the total cohort) | 1.7 (1.0 to 2.0) | Respiratory involvement        | 61 (35.1)  | 18 (17 to 18) |
|                     |                        | Severe respiratory outcomes    | 27 (15.5)  | 18 (17 to 19) |
|                     |                        | Cardiomyopathy                 | 95 (54.6)  | 18 (17 to 18) |

Sensitivity analysis variation #2: One-year observation window; no minimum follow-up imposed

|                      |                        | LOA                            | 86 (33.2) | 10 (9 to 10)  |
|                      |                        | Scoliosis                      | 53 (20.5)  | 10 (9 to 10)  |
|                      |                        | Neurologic/neuropsychiatric involvement | 81 (31.3)  | 9 (9 to 10)  |
| 8 to 10 years (n = 258, 13.2% of the total cohort) | 1.0 (1.0 to 1.0) | LOA                            | 97 (40.8)  | 13 (12 to 13) |
| 11 to 13 years (n = 238, 12.1% of the total cohort) | 1.0 (1.0 to 1.0) | Scoliosis                      | 81 (34.0)  | 13 (12 to 13) |
| 14 to 16 years (n = 281, 14.3% of the total cohort) | 0.9 (1.0 to 1.0) | Neurologic/neuropsychiatric involvement | 74 (31.1)  | 12 (12 to 13) |
| 17 to 19 years (n = 174, 8.8% of the total cohort) | 1.0 (1.0 to 1.0) | Respiratory involvement        | 99 (35.2)  | 15 (15 to 16) |
|                     |                        | Severe respiratory outcomes    | 45 (16.0)  | 15 (15 to 16) |
|                     |                        | Cardiomyopathy                 | 143 (50.9) | 15 (15 to 16) |
|                     |                        | Respiratory involvement        | 57 (32.8)  | 18 (17 to 18) |
|                     |                        | Severe respiratory outcomes    | 24 (13.8)  | 18 (17 to 18) |
|                     |                        | Cardiomyopathy                 | 94 (54.0)  | 18 (17 to 18) |

*Denominator: all patients within the age-range at cohort entry. Abbreviations: IQR = interquartile range, LOA = loss of ambulation.
the severity of the underlying condition; nor are
the underlying reasons for a physician selecting
a particular diagnostic code available. For example,
whether the use of a specific code indicates
a clinical suspicion of early signs of a complica-
tion versus a severe manifestation, or whether a
medication is used prophylactically or for acute treat-
ment, is not clear. Some outcomes may not be easily
identified in administrative data, particularly cap-
ture of initial symptoms or behaviors. For example,
some of the diagnoses contributing to the neu-
ropsychiatric/neurologic involvement outcome are
challenging to identify using claims datasets [43].
Of necessity, selected outcomes in the current study
were ascertained based on proxy measures within the
claims datasets and the reliability of these is unclear
[44]. For LOA, as a specific diagnostic code is not
available, assessment was based on wheelchair use:
However, identification of a wheelchair code does
not indicate if it is a first purchase or a replacement;
or give information of the frequency of wheelchair
use. Equally, wheelchairs may be acquired via other
sources. In a similar vein, for identifying respiratory
involvement, ventilation use may indicate the start
of respiratory decline or a management strategy for
more severe disease. Finally, across all the outcomes
considered, it is possible that a given encounter was
coded as a visit for ‘DMD’, rather than for the specific
manifestation of interest. Despite these challenges,
these data provide indicators of how patients progress
when managed in a standard clinical practice setting
and have implications for using real-world data to
monitor the clinical burden of patients with DMD
over time.

There are some additional limitations to the anal-
yses that should be noted. A clinically validated case
definition for use in administrative claims databases is
not available and there may have been some misclas-
sification on exposure (e.g. patients with other MDs
including limb-girdle muscular dystrophy or BMD
may have been inadvertently included in the study
cohort and patients with DMD over age 30 may have
been inadvertently excluded). Other methods to help
eliminate other MD cases, for example restricting the
timing of the first observed MD diagnosis to early
childhood, were not implemented because of the rel-
atively short follow-up per patient available to allow
true ascertainment of the first diagnosis, and also to
avoid 1) a prohibitively small sample size for analysis,
and 2) disregarding true cases of DMD who entered
the cohort at older ages (due to the limited follow-up
available within MarketScan). Nevertheless, addi-
tional sensitivity analyses performed to explore the
impact of case definition assumptions showed lit-
tle variability in estimates of study outcomes. In
addition, while the sensitivity analysis restricting to
those <18 years at index led to lower median age
estimates, the direction of variation in the frequency
of key clinical outcomes observed supported the
base case findings on the prevalence of these events.
Wheelchair use was consistently documented across
sensitivity analysis cohorts; and events occurring ear-
lier in life were observed at a higher frequency (while
events occurring later in life were observed at a lower

<table>
<thead>
<tr>
<th>Cumulative age-specific prevalence at the end of each follow-up year</th>
<th>Year 1</th>
<th>Year 2</th>
<th>Year 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>LOA</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age 8 to 10; n = 259</td>
<td>24.9%</td>
<td>40.2%</td>
<td>52.9%</td>
</tr>
<tr>
<td>Age 11 to 13; n = 237</td>
<td>37.8%</td>
<td>50.8%</td>
<td>59.0%</td>
</tr>
<tr>
<td>Scoliosis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age 8 to 10; n = 259</td>
<td>15.0%</td>
<td>24.1%</td>
<td>30.4%</td>
</tr>
<tr>
<td>Age 11 to 13; n = 237</td>
<td>29.5%</td>
<td>43.1%</td>
<td>47.4%</td>
</tr>
<tr>
<td>Neurologic/neuropsychiatric involvement</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age 8 to 10; n = 259</td>
<td>32.8%</td>
<td>44.4%</td>
<td>46.1%</td>
</tr>
<tr>
<td>Age 11 to 13; n = 237</td>
<td>26.3%</td>
<td>32.4%</td>
<td>38.0%</td>
</tr>
<tr>
<td>Respiratory involvement</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age 14 to 16; n = 264</td>
<td>27.9%</td>
<td>39.5%</td>
<td>44.4%</td>
</tr>
<tr>
<td>Age 17 to 19; n = 267</td>
<td>35.8%</td>
<td>42.1%</td>
<td>43.5%</td>
</tr>
<tr>
<td>Cardiomyopathy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age 14 to 16; n = 264</td>
<td>44.9%</td>
<td>52.7%</td>
<td>61.3%</td>
</tr>
<tr>
<td>Age 17 to 19; n = 267</td>
<td>54.4%</td>
<td>58.1%</td>
<td>64.1%</td>
</tr>
</tbody>
</table>

Abbreviations: LOA = loss of ambulation.
frequency) among the cohort restricted to those <18 years at index. One additional limitation to note with regard to the definition of DMD, is that individuals with DMD who appear in the MarketScan databases over age 30 years for the first time would be missed; however, this would likely occur quite rarely.

Given the limited follow-up available per patient and that the age at cohort entry of the overall population was in the teenage years, the timing of key clinical milestones occurring prior to cohort entry cannot be accurately ascertained. It therefore cannot be determined whether the first event observed was the first occurrence of the key milestone in a patient's life (rather than an acknowledgement of the patient having already passed that milestone or ongoing documentation of what has become a chronic condition).

As exemplified through the age-restricted sensitivity analyses varying follow-up and observation window requirements, follow-up duration plays a large role in the capture of key milestones in a patient's life. As a result, this study presents the prevalence of key milestones by age at cohort entry and fixed observation windows to allow better interpretation of findings. To supplement this, the KM analyses showed that the prevalence of events captured increases as follow-up increases.

Additionally, a challenge in comparing the occurrence of outcomes with earlier studies is introduced by developments in research and diagnostics such that certain complications (e.g. early signs of cardiomyopathy) may be identified at a younger age through the increased sensitivity of tools such as cardiac MRI. This may therefore lead to apparent higher rates and earlier identification compared to earlier studies with outcomes based on older technology. While understanding mortality is of interest, very limited data were available (i.e. only from inpatient records, and only from the beginning of the study period through 2016); this outcome was not investigated further.

Assessing corticosteroid treatment patterns was not a primary objective of the study and as such it was not designed to comprehensively assess these. As with the analyses of key clinical milestones, it is important to consider the window of follow-up data availability when interpreting available data on corticosteroid use [45]. Given that the majority of this cohort was outside of the age range with the highest anticipated corticosteroid use (6 to 12 years of age), and a large proportion of patients had lost ambulation during the study, the lower rate of overall corticosteroid use in this study cohort is expected and should be interpreted in this context. Finally, the results of this study are specific to individuals covered under a subset of commercial plans and may not be generalizable to all commercial or to government payer segments in the US.

The use of well-validated datasets that provided a large sample size of DMD patients, including children and young adults, from varied commercial insurance plans, was a key strength of this study. Performing age- and time-restricted analyses helped to address potential biases associated with estimating time-dependent outcomes from individuals of differing ages over the course of a non-standardized follow-up window. However, it is important to remember that there is between-patient heterogeneity in the timing of the occurrence of these outcomes, [14, 16] which complicates accurate ascertainment when considered in the context of relatively short follow-up windows per patient available in claims datasets. The results of the sensitivity analyses for the cohort definition were consistent with those of the primary analyses, providing support that the study findings were robust. With respect to the cohort definition, identifying patients with DMD-genotype specific treatments (such as eteplirsen) or DMD/BMD-specific ICD-10 codes (G71.01) was also explored, but did not identify any additional patients beyond those already captured by the study inclusion criteria.

**CONCLUSIONS**

This is the first study to document the occurrence of key clinical milestones among real-world patients with DMD with a wide range of commercial insurance plans. The prevalence of key clinical milestones observed by age was broadly consistent with published findings from large registries. Variability in estimates reflect the clinical heterogeneity experienced by those with DMD but also the impact of observation windows on the ability to ascertain key clinical outcomes in DMD. As such, the findings of this study also help delineate the types of outcomes that one can well characterize using existing claims data for cohort studies, and considerations on the methodology to do so. These data summarizing the occurrence of relevant clinical outcomes among patients with DMD with commercial insurance coverage add to the growing body of evidence describing the clinical course of DMD patients using real-world data.
REFERENCES


[23] IBM Watson Health. COMMERCIAL CLAIMS AND ENCOUNTERS DATABASE AND MEDICARE
SUPPLEMENTAL AND COORDINATION OF BENEFITS DATABASE. IBM Watson Health; 2017.


[38] 38.. !!! INVALID CITATION !!! 16.


