Response to treatment of myasthenia gravis according to clinical subtype

Tetsuya Akaishi¹, Yasushi Suzuki², Tomihiro Imai³, Emiko Tsuda³, Naoya Minami⁴, Yuriko Nagane⁵, Akiyuki Uzawa⁶, Naoki Kawaguchi⁶, Masayuki Masuda⁷, Shingo Konno⁸, Hidekazu Suzuki⁹, Hiroyuki Murai¹⁰, Masashi Aoki¹ and Kimiaki Utsugisawa⁵*

Abstract

Background: We have previously reported using two-step cluster analysis to classify myasthenia gravis (MG) patients into the following five subtypes: ocular MG; thymoma-associated MG; MG with thymic hyperplasia; anti-acetylcholine receptor antibody (AChR-Ab)-negative MG; and AChR-Ab-positive MG without thymic abnormalities. The objectives of the present study were to examine the reproducibility of this five-subtype classification using a new data set of MG patients and to identify additional characteristics of these subtypes, particularly in regard to response to treatment.

Methods: A total of 923 consecutive MG patients underwent two-step cluster analysis for the classification of subtypes. The variables used for classification were sex, age of onset, disease duration, presence of thymoma or thymic hyperplasia, positivity for AChR-Ab or anti–muscle-specific tyrosine kinase antibody, positivity for other concurrent autoantibodies, and disease condition at worst and current. The period from the start of treatment until the achievement of minimal manifestation status (early-stage response) was determined and then compared between subtypes using Kaplan-Meier analysis and the log-rank test. In addition, between subtypes, the rate of the number of patients who maintained minimal manifestations during the study period/that of patients who only achieved the status once (stability of improved status) was compared.

Results: As a result of two-step cluster analysis, 923 MG patients were classified into five subtypes as follows: ocular MG (AChR-Ab-positivity, 77%; histogram of onset age, skewed to older age); thymoma-associated MG (100%; normal distribution); MG with thymic hyperplasia (89%; skewed to younger age); AChR-Ab-negative MG (0%; normal distribution); and AChR-Ab-positive MG without thymic abnormalities (100%, skewed to older age). Furthermore, patients classified as ocular MG showed the best early-stage response to treatment and stability of improved status, followed by those classified as thymoma-associated MG and AChR-Ab-positive MG without thymic abnormalities; by contrast, those classified as AChR-Ab-negative MG showed the worst early-stage response to treatment and stability of improved status.

Conclusions: Differences were seen between the five subtypes in demographic characteristics, clinical severity, and therapeutic response. Our five-subtype classification approach would be beneficial not only to elucidate disease subtypes, but also to plan treatment strategies for individual MG patients.

Keywords: Classification, Cluster analysis, Myasthenia, Onset age, Treatment

* Correspondence: kutsugi@k-dion.ne.jp

© The Author(s). 2016 Open Access This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.
Background

Myasthenia gravis (MG) is a neurological disorder that manifests as fatigable and fluctuating weakness of voluntary muscles, which are mediated by autoantibodies against neuromuscular junction proteins in skeletal muscle that impair neuromuscular transmission [1]. MG typically involves the ocular, bulbar, and extremity muscles, and, in severe cases, respiratory muscles. The clinical course and outcome in MG are affected by several different autoantibodies, thymic abnormalities, onset age and disease severity, as well as response to treatment [2–4]. MG is distinguished according to the production of pathogenic autoantibodies such as anti-acetylcholine receptor antibody (AChR-Ab) and anti–muscle-specific tyrosine kinase antibody (MuSK-Ab) [1, 5, 6]. Clinically, MG is often classified into the following three subtypes based on thymic abnormalities and onset age: thymoma-associated MG; early-onset MG (onset age <50 years); and late-onset MG (onset age ≥50 years) [7]. Furthermore, discrimination is observed in the clinical setting—for example, between ocular and generalized MG—based on the distribution of symptoms.

Previously, we reported classifying MG into the following five subtypes using two-step cluster analysis of a detailed cross-sectional data set of 640 consecutive patients (Japan MG Registry Study 2012): ocular MG; generalized thymoma-associated MG; generalized MG with thymic hyperplasia; generalized AChR-Ab-negative MG; and generalized AChR-Ab-positive MG without thymic abnormalities [8]. However, this five-subtype classification approach requires further confirmation, and its clinical relevance remains to be established.

Therefore, in 2015, we conducted a larger cross-sectional survey to obtain clinical parameters from 1,088 consecutive MG patients. In the present study, using this new data set, we attempted to confirm the reproducibility of our five-subtype classification approach and to specify additional characteristics of these five subtypes with a particular focus on response to treatment in the clinical setting.

Methods

Patients and clinical factors

This survey was conducted by the Japan MG Registry Study Group, which comprises 13 neurological centers (Table 1). We evaluated patients with established MG between April and July 2015. To avoid potential bias, we enrolled consecutive patients over a short duration (4 months). All 1088 of these MG patients visited our hospitals, provided written informed consent, and underwent analysis. Among these 1088 patients, 331 (30.4%) were included in our previous survey in 2012 [8].

Table 1 Institutions participating in the Japan MG Registry Study 2015

| Institution                                                                 |
|--------------------------------------------------------------------------------|
| Department of Neurology, Sapporo Medical University Hospital, Sapporo        |
| Department of Neurology, Hokkaido Medical Center, Sapporo                    |
| Department of Neurology, Hanamaki General Hospital, Hanamaki                |
| Department of Neurology, Sendai Medical Center, Sendai                       |
| Department of Neurology, Tohoku University Graduate School of Medicine, Sendai |
| Chiba Neurology Clinic, Chiba                                                |
| Department of Neurology, Chiba University School of Medicine, Chiba          |
| Department of Neurology, Tokyo Medical University, Tokyo                     |
| Department of Neurology, Toho University Medical Center Oh-hashi Hospital, Tokyo |
| Department of Neurology, Tokyo Women’s Medical University, Tokyo             |
| Department of Neurology, Kinki University School of Medicine, Osaka          |
| Department of Neurology, Graduate School of Medical Sciences, Kyushu University, Fukuoka |
| Department of Neurology and Strokology, Nagasaki University Hospital, Nagasaki |

Abbreviation: MG myasthenia gravis

The following clinical parameters were obtained for all patients: sex; age; age at disease onset; duration of disease; duration of immunotherapy; history of bulbar symptoms; presence of thymoma or thymic hyperplasia in thymectomized patients; presence of serum AChR-Ab or MuSK-Ab; and presence of other non-MG-specific autoantibodies, such as anti-nuclear antibody, SS-A/SS-B antibody, TSH-receptor antibody, anti-thyroglobulin/thyroxoperoxidase antibody, and rheumatoid factor. In addition, the current and past disease status and details of treatment were surveyed for all patients. Clinical severity at the worst condition was determined according to the classification of the MG Foundation of America (MGFA) [9], and, in some patients, the MGFA quantitative MG score (QMG) [9, 10] from medical records. Clinical severity at the current condition was determined according to QMG and MG Composite (MGC) scores [11]. Furthermore, all patients completed the Japanese version of the 15-item Myasthenia Gravis Quality-of-Life Scale (MG-QOL-15), [12, 13] upon study entry.

Prednisone and prednisolone are the global standard oral corticosteroids used to treat MG, and prednisolone is generally used in Japan. Therefore, the current use, peak dose [mg/day], and duration of prednisolone ≥20 mg/day were recorded for all patients, as was the use of calcineurin inhibitors, azathioprine, plasmapheresis, and intravenous immunoglobulin.

Finally, the courses of current and past MGFA post-intervention statuses, particularly the time required to achieve first minimal manifestations (MM) or better status lasting more than one month (MM-or-better ≥1 M) [9], were determined as benchmarks for evaluating
response to treatment in each patient. These clinical data were fully collected from 923 (84.8%) of the 1088 patients.

Two-step cluster analysis
To examine the reproducibility of the five-subtype classification in the same manner as reported elsewhere [8], we conducted two-step cluster analysis of the 923 patients using SPSS Statistics Base 22 software (IBM, Armonk, NY, USA). To avoid bias beset by the problem of multicollinearity, current or worst disease status was handled as a single variable (Table 2). The other variables evaluated were: sex; age of onset; disease duration; presence of thymoma; presence of thymic hyperplasia in thymectomized cases; positivity for AChR-Ab or MuSK-Ab; and positivity for other concurrent autoantibodies (Table 2).

Early-stage response to treatment and stability of improved status in each of the five subtypes

Early-stage response to treatment
The time (months) from the start of the immunotherapy until achieving first MM-or-better ≥1 M was determined from medical records and compared between the five subtypes using Kaplan-Meier analysis and the log-rank test with the Cochran-Mantel-Haenszel procedure. The time required to achieve first MM-or-better ≥1 M in 50% of patients was also compared among subtypes.

Stability of improved status of MM-or-better ≥1 M
As an indicator of stability of improved status, the rate of the number of patients who maintained minimal manifestations in the 2015 survey/that of patients who achieved the status at least once was calculated and compared among the five subtypes.

Statistical analysis
All statistical analyses were performed using SPSS Statistics Base 22 software (IBM) and MATLAB R2015a (MathWorks, Natick, MA, USA). All continuous data are expressed as the mean ± standard deviation (SD) and the median.

Results

Two-step cluster analysis
Based on the results of two-step cluster analyses, all 923 MG patients could be classified into the same five subtypes described elsewhere [8]: ocular MG; thymoma-associated MG; MG with thymic hyperplasia; AChR-Ab-negative MG; and other (in order of predicted importance). Among these five subtypes, the residual patients group “other” was the largest, and could be defined as generalized AChR-Ab-positive MG without thymic abnormalities. These results were demonstrated repeatedly with several sets of variables, as shown in Table 2, which confirmed the high reliability and reproducibility of the classification system. Although the order among thymoma-associated MG, MG with thymic hyperplasia, and AChR-Ab-negative MG was unstable depending on the variable sets used, the differences in terms of predicted importance were not large. These results were almost identical to those reported elsewhere [8], with only minor discrepancies in regard to the order of selection priority (the order in the previous study was as follows: ocular MG; MG with thymic hyperplasia; AChR-Ab-negative MG; thymoma-associated MG; and AChR-Ab-positive MG without thymic abnormalities). In the present study, the quality of clusterization under each set of variables, which was estimated using a previously reported interpretation model [14], was indicated as “fair” to “good” for all clusters, suggesting that the results were reasonable.

A total of 111 patients (10.2%) fit two of the five subtypes (Table 3). These patients were allocated to sole subtypes according to the separation priority in the two-step cluster analysis. For example, an ocular MG patient with thymoma was allocated into ocular MG. Under this criterion, the percentage of patients assigned to the five subtypes was as follows: ocular MG, 23.0%; thymoma-associated MG, 21.5%; MG with thymic hyperplasia, 12.9%; AChR-Ab-negative MG, 12.1%; and AChR-Ab-positive MG without thymic abnormalities, 30.5% (Table 4).

MG with thymic hyperplasia is only diagnosed for thymectomized patients; therefore, some non-thymectomized patients with thymic hyperplasia may be assigned as other subtypes, particularly AChR-Ab-positive MG patients without thymic abnormalities.

| Patients’ backgrounds | Autoantibody status | Disease status during the worst condition | Current disease status |
|-----------------------|---------------------|------------------------------------------|-----------------------|
| Sex                   | AChR-Ab             | One of the following:                   | One of the following: |
| Age of onset          | MuSK-Ab             | The worst MGFA classification           | Current PIS           |
| Disease duration      | Non-MG-specific antibodies | The worst QMG                        | Current QMG           |
| Presence of thymoma   |                     |                                          | MG-QOL-15 score       |
| Presence of thymic hyperplasia |                 |                                          |                       |

AChR-Ab anti-acetylcholine receptor antibody, MuSK-Ab anti-muscle specific kinase antibody, MG myasthenia gravis, MGFA MG Foundation of America, QMG quantitative MG, MG-QOL-15 15-item MG-specific quality of life scale, PIS post-intervention status
The clinical characteristics, including current and worst severity, for each of the five subtypes are shown in Table 4. The patients with MuSK-Ab were not separated by two-step cluster analysis because the number is not great enough for statistical evaluation. However, MG patients with MuSK-Ab showed a distinct clinical manifestation and therapy responsiveness reflecting the unique pathological mechanism [15]. Therefore, details of MG patients with MuSK-Ab (n = 22) are individually described next to SNMG patients in Table 4. The percentage of females was significantly higher among MG patients with MuSK-Ab (n = 22) compared to other subtypes. The worst condition of the disease was MGC score (mean ± SD, median), which was significantly higher in the MuSK-Ab-positive group and MGFA > III% in MG patients with MuSK-Ab, compared to other subtypes. The QMG score (n = 225), MG-QOL-15 (n = 286), and MGC score (n = 286) were also assessed, and the results were comparable to those of the MuSK-Ab-positive group. All continuous data are expressed as the mean ± standard deviation (SD) and the median. *p < 0.0001, chi-square test (compared to the others), †p < 0.0001, Mann–Whitney U test.

### Table 3 Number of patients fitting two categories

|                  | Number of patients | Final assignment |
|------------------|--------------------|-----------------|
| Ocular MG and thymoma-associated MG | 32 (2.9%) | Ocular MG |
| Ocular MG and AChR-Ab-negative MG | 56 (5.1%) | Ocular MG |
| Ocular MG and MG with thymic hyperplasia | 8 (0.7%) | Ocular MG |
| MG with thymic hyperplasia and AChR-Ab-negative MG | 14 (1.3%) | THMG |
| Thymoma-associated MG and AChR-Ab-negative MG | 1 (0.09%) | TAMG |

Values within the parentheses show the percentages of the total of 1,088 patients.

### Table 4 Characteristics and severity for each of the five MG subtypes

|                  | Ocular MG | Thymoma-associated MG | MG with thymic hyperplasia | AChR-Ab-negative MG | (MuSK-Ab-positive) | AChR-Ab-positive MG without thymic abnormalities | Total |
|------------------|-----------|-----------------------|---------------------------|---------------------|-------------------|-----------------------------------------------|-------|
| Patients (n)     | 250       | 234                   | 140                       | 132 (22)            | 332               | 1088                                          |       |
| Female,%         | 52.0      | 67.5                  | 81.4*                     | 81.1* (81.8)        | 61.1              | 65.4                                          |       |
| Onset age, y     | 51.0 ± 20.0, 53.0* | 51.0 ± 12.8, 51.0      | 33.3 ± 13.9, 31.0*        | 39.9 ± 16.2, 41.5*  | 50.7 ± 20.7, 55.0* | 47.3 ± 18.8, 48.1                                  |       |
| Duration of disease, y | 10.4 ± 11.4, 6.4 | 9.5 ± 7.7, 8.0          | 17.4 ± 11.8, 14.5*        | 10.8 ± 9.3, 8.0     | 11.6 ± 10.5, 8.2 | 11.6 ± 10.6, 8.2                                  |       |
| AChR-Ab-positivity,% | 77.2      | 99.6                  | 89.4                      | 0.0                 | 100.0             | 81.3                                          |       |
| MuSK-Ab-positivity,% | 0.0%      | 0.0%                  | 0.0%                      | 20.6%               | 0.0%              | 2.1%                                          |       |
| Thymectomy,%     | 23.6%*    | 97.4%*                | 100.0%*                   | 12.1%*              | 35.8%*            | 51.7%                                         |       |

The worst condition of the disease

- **MGFA classification (n = 1088):**
  - I,% 100.0%
  - II,% 0.0%
  - III,% 0.0%
  - IV,% 0.0%
  - V,% 0.0%

- **Rate of MGFA > III%:**
  - 0.0%
  - 56.0%*

- **QMG score (n = 922):**
  - 6.6 ± 2.6, 6.0 (n = 225)
  - 17.1 ± 8.0, 15.5 (n = 194)
  - 15.8 ± 5.8, 15.0 (n = 107)
  - 14.7 ± 7.2, 13.0 (n = 114)
  - 18.1 ± 9.7, 15.5 (n = 107)
  - 14.7 ± 7.0, 13.0 (n = 114)
  - 13.4 ± 7.5, 12.0 (n = 114)

- **MGC score (n = 923):**
  - 4.2 ± 2.8, 4.0 (n = 208)
  - 6.8 ± 4.8, 6.0 (n = 198)
  - 7.8 ± 5.5, 7.0 (n = 125)
  - 8.4 ± 5.4, 8.0 (n = 106)
  - 8.3 ± 6.4, 7.0 (n = 106)
  - 7.2 ± 4.8, 6.0 (n = 106)
  - 6.6 ± 4.8, 6.0 (n = 106)

- **MG-QOL-15 (n = 923):**
  - 8.1 ± 9.0, 5.0 (n = 208)
  - 14.7 ± 13.6, 12.0 (n = 125)
  - 16.2 ± 13.7, 13.5 (n = 106)
  - 14.6 ± 12.6, 11.0 (n = 106)
  - 11.6 ± 8.8, 11.0 (n = 106)
  - 14.1 ± 13.9, 9.0 (n = 106)
  - 13.2 ± 13.0, 9.0 (n = 106)

All continuous data are expressed as the mean ± standard deviation (SD) and the median. *p < 0.0001, chi-square test (compared to the others), †p < 0.0001, Mann–Whitney U test.
with thymic hyperplasia and AChR-Ab-negative MG patients compared with the other three subtypes \((p < 0.0001, \text{chi-square test})\). Onset age was significantly younger in MG with thymic hyperplasia and AChR-Ab-negative MG patients \((p < 0.0001, \text{Mann–Whitney U test})\) and older in ocular MG and AChR-Ab-positive MG patients without thymic abnormalities \((p < 0.0001, \text{Mann–Whitney U test})\).

Severity at the worst condition (MGFA classification and QMG) was significantly higher in thymoma-associated MG patients \((p < 0.0001, \text{Mann–Whitney U test})\). Patients with MuSK-Ab also showed worst severity at the same level as thymoma-associated MG patients, although this result was not statistically significant because of the small number of patients. The severity scales (QMG and MGC), and a QOL scale (MG-QOL-15) scores in the present survey were generally worse, although not statistically significant, in MG with thymic hyperplasia and AChR-Ab-negative MG patients, both of which primarily comprise females with younger onset ages. On the other hand, as a matter of course, ocular MG patients showed much lower clinical severity in all batteries at both current and worst conditions \((p < 0.0001, \text{Mann–Whitney U test})\).

**Onset age histograms of the five subtypes**

Histograms of the onset age for each of the five subtypes are shown in Fig. 1a. These histograms were converted into approximate curves (sixth-order polynomial approximations) and superimposed in Fig. 1b. The peak ages of the histogram were 60–64 years in ocular MG, 25–29 years in MG with thymic hyperplasia, 35–39 years in AChR-Ab-negative MG, 50–54 years in thymoma-associated MG, and 65–69 years in AChR-Ab-positive MG without thymic abnormalities. The histogram was skewed toward younger onset age in MG with thymic hyperplasia and toward older onset age in ocular MG and AChR-Ab-positive MG without thymic abnormalities. The histogram was skewed toward younger onset age in MG with thymic hyperplasia and toward older onset age in ocular MG and AChR-Ab-positive MG without thymic abnormalities. Regarding patients with MuSK-Ab \((n = 22)\), the mean \(±\) SD onset age was 38.6 \(±\) 15.3 (median, 42.0 years), and the ratio of females was 81.8%; however, neither of these findings was significantly different from other AChR-Ab-negative MG patients.

**Early-stage response to treatment and stability of improved status among the five subtypes**

Details of past immunotherapy for each of the five subtypes are shown at the top of Table 5.

**Early-stage response to treatment (first achievement of MM-or-better ≥1 M)**

As shown in the middle of Table 5, the rate of the patients achieving MM-or-better ≥1 M at least once was significantly higher for ocular MG \((p < 0.001, \text{chi-square test})\) and significantly lower for AChR-Ab-negative MG \((p < 0.001, \text{chi-square test})\).

Kaplan-Meier curves for the time to first achieve MM-or-better ≥1 M in each of the five subtypes up to 10 years from initiating immunotherapy are shown in Fig. 2a. The time to first achieve MM-or-better ≥1 M was significantly different among the five subtypes \((p < 0.0001; \text{generalized Wilcoxon test and log-rank test})\). Significant differences were observed between all pairs of two subtypes \((p < 0.01\) for all pairs; generalized Wilcoxon test) except MG with thymic hyperplasia and AChR-Ab-negative MG \((p ≥ 0.10)\). Patients with ocular MG showed the best early-stage response to treatment compared with others \((p < 0.0001; \text{log-rank test}, p < 0.001; \text{chi-square test})\). The time required to achieve MM-or-better ≥1 M in 50% of the patients was significantly longer in MG with thymic hyperplasia and AChR-Ab-negative MG compared with ocular MG, thymoma-associated MG, and AChR-Ab-positive MG without thymic abnormalities \((p < 0.0001, \text{log-rank test}; \text{middle of Table 5})\). For comparison, Kaplan-Meier curves for the time to first achieve MM-or-better ≥1 M in early-onset, late-onset, and thymoma-associated MG (three-type classification) are shown in Fig. 2b. Significant differences were observed between early- and late-onset MG \((p < 0.01)\) and between early-onset and thymoma-associated MG \((p < 0.01)\); however, no significant differences were found between late-onset and thymoma-associated MG \((p ≥ 0.10)\).

**Stability of improved status**

The rates of patients with MM-or-better status during the survey and stability of improved status are shown in the bottom of Table 5. Stability of improved status was significantly better in ocular MG compared with other subtypes \((p < 0.0001; \text{chi-square test})\); however, no significant differences were observed among subtypes other than ocular MG \((p ≥ 0.10\) for all pairs, excluding ocular MG; chi-square test).

**Discussion**

The present analyses based on several sets of variables classified 923 MG patients into the same five following subtypes with the same characteristics of the onset-age histograms as reported in our previous study [8]: ocular MG (AChR-Ab-positivity, 77%; histogram of onset age, skewed to older age); thymoma-associated MG (100%; normal distribution); MG with thymic hyperplasia (89%; skewed to younger age); AChR-Ab-negative MG (0%; normal distribution); and AChR-Ab-positive MG without thymic abnormalities (100%, skewed to older age). The results from the two different samples demonstrated high reproducibility, which suggests the reliability of our five-subtype classification method. In the process of
Fig. 1 (See legend on next page.)
analyses, two points were suggested. First, discrimination between ocular and generalized MG is more principal than that according to onset age, thymus pathology or AChR-Ab-positivity. Second, AChR-Ab-negative MG shows normal distribution of onset age not fitting discrimination based on onset age. Therefore, it is probably better to adopt the often-used three-type classification (early-onset, late-onset and thymoma-associated MG) for generalized and AChR-Ab-positive phenotypes. Consistently, in our five-subtype classification, MG with thymic hyperplasia (with early-onset age), AChR-Ab-positive MG without thymic abnormalities (with late-onset age) and thymoma-associated MG were generalized and AChR-Ab-positive phenotypes.

In fact, these results of our classification statistically performed are consistent with a recently reported classification of MG by Gilhus et al. [16, 17], which included the following classifications: early-onset MG; late-onset MG; thymoma-associated MG; MuSK-Ab positive MG; lipoprotein-related protein 4 (LRP4)-Ab positive MG; seronegative MG; and ocular MG. In addition, they commented that early- and late-onset MG should be distinguished according to onset age only for patients having generalized symptoms and AChR-Ab. In the present study, because of their small numbers, MuSK-Ab-positive MG patients were not separated, and LRP4-Ab positivity was not systematically determined; though MG patients with MuSK-Ab or LRP 4-Ab have a distinct clinical manifestation and a unique pathological mechanism [15].

Ocular MG was found to have unique characteristics such as having a higher onset age, predominantly affecting males, and having an ocular muscle-specific pathogenesis [18], which may be related to the aging-associated susceptibility of ocular muscles to antibodies against the neuromuscular junction. Given that response to treatment and stability of improved status were substantially better in ocular MG compared with the other four subtypes, it seems reasonable to conclude that ocular MG should be treated as a distinct subgroup of MG in the clinical setting.

Among the four generalized subtypes in the present classification method, both early-stage response to treatment and stability of improved status were worst in

**Table 5** Details of past treatment and response to treatment for each of the five subtypes

| Past immunotherapy (n = 923) | Ocular MG | Thymoma-associated MG | MG with thymic hyperplasia | AChR-Ab-negative MG | (MuSK-Ab-positive) | AChR-Ab-positive MG without thymic abnormalities | Total |
|------------------------------|-----------|------------------------|----------------------------|---------------------|-------------------|-----------------------------------------------|-------|
| Peak dose of oral PSL, mg/day | 9.2 ± 12.2, 5.0⁺ | 28.5 ± 18.8, 30.0⁺ | 29.7 ± 19.4, 30.0⁺ | 18.8 ± 17.2, 15.0 | 32.6 ± 20.6, 30.0 | 23.7 ± 20.2, 20.0 | 21.5 ± 19.3, 15.0 |
| Duration of PSL ≥20 mg/day, M | 0.0 ± 0.0, 0.0⁺ | 12.0 ± 25.2, 5.0⁺ | 13.0 ± 27.3, 6.0⁺ | 3.8 ± 7.0, 0.0 | 7.2 ± 9.5, 4.0 | 8.2 ± 17.0, 2.0 | 7.9 ± 19.3, 1.0 |
| CNIs,% | 24.0%* | 68.2%* | 54.0% | 67.4% | (72.7%) | 58.1% | 52.9% |
| PP,% | 2.0%* | 48.1%* | 22.1% | 46.0% | (54.5%) | 37.2% | 27.3% |
| IVIG,% | 6.1%* | 36.1% | 29.9% | 42.5% | (27.3%) | 24.7% | 15.0% |
| Initial response to treatment (n = 923, see Fig. 2a) | Achievement of MM-or-better once,% | 79.8%* | 73.5% | 66.1% | 56.2%* | (75.0%) | 67.8% | 70.2% |
| Months to achieve MM-or-better in 50% of patients | 4.0⁺ | 8.0 | 18.0⁺ | 31.0⁺ | (7.0) | 6.0 | 8.0 |
| Stability of improved status (n = 923) | MM-or-better at present,% | 74.0%* | 58.1% | 49.6% | 39.6%* | (55.0%) | 55.4% | 57.6% |
| Maintaining rate of MM-or-better,% | 92.7%* | 79.0% | 75.0% | 70.5% | (73.3%) | 81.7% | 82.1% |

*All continuous data are expressed as the mean ± standard deviation (SD) and the median
CNIs calcineurin inhibitors, EAT early aggressive therapy, IVIG intravenous immunoglobulin, M months, MG myasthenia gravis, MM-or-better ≥1 M minimal manifestation or better status lasting more than one month, MuSK-Ab-positive MG patients with serum anti-muscle specific kinase (MuSK) autoantibody in AChR-Ab-negative patients, PP plasmapheresis, PSL prednisolone, SD standard deviation
*p < 0.0001, chi-square test, †p < 0.0001, Mann-Whitney U test, ‡p < 0.0001, log-rank test
AChR-Ab-negative MG, although symptoms at the time the condition was at its worst were not particularly severe. Patients with MuSK-Ab-positive MG showed better results despite having more severe worst conditions, which suggests that AChR-Ab-negative MG, excluding MuSK-Ab-positive MG, is distinct from other generalized MG subtypes from the perspective of response to therapy. Overall, as shown in Fig. 2a, each of the five present subtypes showed different levels of response to treatment, whereas such differences in the three commonly used subtypes (early-onset, late-onset, and thymoma-associated MG) remain somewhat unclear (Fig. 2b). It would be more helpful in the clinical setting to elucidate the levels of response to some types of
medication or therapy (e.g. corticosteroids, non-steroid immunosuppressants, intravenous immunoglobulin and plasmapheresis) in the five subtypes. However, it was difficult to analyze such response levels, as plural treatment agents and methods were employed simultaneously in most of individual patients. We are now analyzing the response levels according to patterns of immune treatment (treatment strategies) in generalized MG patients [19]. Such analysis should be performed also for the present five subtypes, but could not be addressed in the present report.

The present study did have some limitations. First, 331 (30.4%) of the 1,088 patients were included in our previous survey in 2012, which might have affected the reproducibility of the present five-subtype classification. Second, almost all of the MG patients in our database are Japanese; therefore, a race/ethnicity bias may have affected the results. Finally, some MG patients with thymic hyperplasia might have been classified as AChR-Ab-positive MG without thymic abnormalities because the diagnosis of thymic hyperplasia is made based on the results of pathological examinations after thymectomy. However, the frequency of thymectomy for MG patients without thymoma has been decreasing [20]; therefore, some of the AChR-Ab-positive MG patients with younger onset age who had not undergone thymectomy could have had thymic hyperplasia.

Conclusion

The results of the present study suggest that MG patients can be classified into the following five subtypes in order of priority: ocular MG; thymoma-associated MG; MG with thymic hyperplasia; AChR-Ab-negative MG; and AChR-Ab-positive MG without thymic abnormalities. All MG patients can be allocated to one of the subtypes based on the results of routine examinations. These five subtypes were shown to have characteristic demographic characteristics, clinical severity, and therapeutic responses. Therefore, our five-subtype classification method is expected to be beneficial not only for elucidating disease types, but also for planning proper treatment for individual patients.

Acknowledgements

The authors would like to thank Dr. M. Motomura and Dr. H. Shiraishi (Department of Neurology and Strokeology, Nagasaki University Hospital), Dr. Y. Shimizu and Dr. R. Ikeguchi (Department of Neurology, Tokyo Women’s Medical University), Dr. D. Yamamoto (Department of Neurology, Sapporo Medical University Hospital) and Dr. T. Kanai (Department of Neurology, Chiba University School of Medicine) for collecting the patient data.

Funding

This work was supported by the Japan MG Registry study group.

Availability of data and material

The data sets during and/or analyzed during the current study are available from the corresponding author on reasonable request.

Authors’ contributions

Study concept and design: TA, YS, HM, MA, KU. Acquisition of data, or analysis and interpretation of data: TA, YS, TI, ET, NM, YM, YN, AU, NK, MM, SK, HS, HM, MA, KU. Drafting the article or revising it critically for important intellectual content: TA, YS, TI, ET, NM, YM, YN, AU, NK, MM, SK, HS, HM, MA, KU. Final approval of the version to be published; and agreement to be accountable for all aspects of the work: TA, YS, TI, ET, NM, YM, YN, AU, NK, MM, SK, HS, HM, MA, KU. Obtained funding and study supervision: HM, MA, KU.

Competing interests

The authors declare that they have no competing interests.

Consent for publication

not applicable

Ethics approval and consent to participate

The study protocols were approved by the ethics committees of each participating institution. Written informed consent was obtained from all patients participating in the study.

Author details

1Department of Neurology, Tohoku University Graduate School of Medicine, Sendai, Japan. 2Department of Neurology, Sendai Medical Center, Sendai, Japan. 3Department of Neurology, Sapporo Medical University, Sapporo, Japan. 4Department of Neurology, Hokkaido Medical Center, Sapporo, Japan. 5Department of Neurology, Hanamaki General Hospital, 4-28 Kajoh-chou, Hanamaki 025-0075, Japan. 6Department of Neurology, Chiba University Graduate School of Medicine, Chiba, Japan. 7Department of Neurology, Tokyo Medical University, Tokyo, Japan. 8Department of Neurology, Toho University School of Medicine, Tokyo, Japan. 9Department of Neurology, Kinki University School of Medicine, Osaka, Japan. 10Department of Neurological Therapeutics, Graduate School of Medical Sciences, Kyushu University, Fukuoka, Japan.

Received: 25 August 2016 Accepted: 11 November 2016

Published online: 17 November 2016

References

1. Vincent A, Leite MI. Neuromuscular junction autoimmune disease: muscle specific kinase antibodies and treatments for myasthenia gravis. Curr Opin Neurol. 2000;13:159–25.
2. Meriggioli MN, Sanders DB. Autoimmune myasthenia gravis: emerging clinical and biological heterogeneity. Lancet Neurol. 2009;8:475–90.
3. Evoli A, Batocchi AP, Minisci C, Di Schino C, Tonnali P. Clinical characteristics and prognosis of myasthenia gravis in older people. J Am Geriatr Soc. 2000;48:1442–8.
4. Grob D, Arusa EL, Brunner NG, Namba T. The course of myasthenia gravis and therapies affecting outcome. Ann N Y Acad Sci. 1987;505:472–99.
5. Lavnic D, Losen M, Vujic A, De Baets M, Hajdukovic LJ, Stojanovic V, et al. The features of myasthenia gravis with autoantibodies to MuSK. J Neurol Neurosurg Psychiatry. 2005;76:1099–102.
6. Gilhus NE. Autoimmune myasthenia gravis. Expert Rev Neurother. 2009;9:351–8.
7. Aarli JA. Late-onset myasthenia gravis: a changing scene. Arch Neurol. 1999;56:25–7.
8. Akiishi T, Yamaguchi T, Suzuki Y, Nagane Y, Suzuki S, Murali H, et al. Insights into the classification of myasthenia gravis. PLoS One. 2014;9:e106757.
9. Jozefczuk 3rd A, Barohn RJ, Ernstoff RW, Kaminski HJ, Keesey JC, Penn AS, et al. Myasthenia gravis: recommendations for clinical research standards. Task Force of the Medical Scientific Advisory Board of the Myasthenia Gravis Foundation of America. Ann Thorac Surg. 2000;70:327–34.
10. Bedirks S, Simel DL, Bosworth H, Samsa G, Tucker-Lipscomb B, Sanders DB. Quantitative myasthenia gravis score: assessment of responsiveness and longitudinal validity. Neurology. 2005;64:1968–70.
11. Burns TM, Conaway M, Sanders DB. The MG Composite: A valid and reliable outcome measure for myasthenia gravis. Neurology. 2010;74:1434–40.
12. Burns TM, Conaway MR, Cutter GR, Sanders DB. Less is more, or almost as much: a 15-item quality-of-life instrument for myasthenia gravis. Muscle Nerve. 2008;38:557–63.
13. Masuda M, Utsugisawa K, Suzuki S, Nagane Y, Kabasawa C, Suzuki Y, et al. The MG-QOL15 Japanese version: validation and associations with clinical factors. Muscle Nerve. 2012;46:166–73.
14. Rousseeuw PJ. Silhouettes: a graphical aid to the interpretation and validation of cluster analysis. J Comput Appl Math. 1987;20:53–65.
15. Huijbers MG, Zhang W, Klooster R, Nks EH, Friese MB, Staaarshem KR, et al. MuSK IgG4 autoantibodies cause myasthenia gravis by inhibiting binding between MuSK and LR4. Proc Natl Acad Sci U S A. 2013;110:20783–8.
16. Gilhus NE, Verschuuren JJ. Myasthenia gravis: subgroup classification and therapeutic strategies. Lancet Neurol. 2015;14:1023–36.
17. Gilhus NE, Skeie GO, Rorni F, Lazaridis K, Zisimopoulou P, Tzartos S. Myasthenia gravis - autoantibody characteristics and their implications for therapy. Nat Rev Neurol. 2016;12:259–68.
18. Kaminski HJ, Ruff RL, Ocular muscle involvement by myasthenia gravis. Ann Neurol. 1997;41:419–20.
19. Utsugisawa K, Nagane Y, Akaishi T, Suzuki Y, Imai T, Tsuda E, et al. Early fast-acting treatment strategy against generalized myasthenia gravis. Muscle Nerve. 2016; mus.25397. [Epub ahead of print]
20. Murai H. Japanese clinical guidelines for myasthenia gravis: Putting into practice. Clin Exp Neuroimmunol. 2015;6:21–31.