Serological diagnosis of myasthenia gravis and its clinical significance

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Abstract: Myasthenia gravis (MG) is the most common immune-mediated disorder of the neuromuscular junction. Anti-acetylcholine receptor (anti-AChR), anti-muscle-specific kinase (anti-MuSK), and anti-lipoprotein receptor-related protein 4 (anti-LRP4) antibodies are the three well-defined pathogenic antibodies. Patients with MG can also have other antibodies, such as anti-titin, anti-ryanodine receptor (anti-RyR), anti-Agrin and anti-KV1.4 antibodies. Since MG is heterogeneous in terms of pathophysiology, antibody status, and other factors, serological tests are crucial for clinical diagnosis confirmation and treatment choice. Herein, we review the different methods for detection of various antibodies involved in MG and their sensitivity and specificity. The understanding of these elements should be useful for improving the diagnosis and determining how to adapt the existing therapies to the requirements of each patient.

Keywords: Myasthenia gravis (MG); antibodies; radioimmunoprecipitation assay (RIPA); enzyme-linked immunosorbent assay (ELISA); cell-based assay (CBA)

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Background

Myasthenia gravis (MG) is an autoimmune disorder caused by antibodies targeting the neuromuscular junction, which has a reported worldwide prevalence of 40–180 per million people, and an annual incidence of 4–12 per million people (1-4). The antibodies attack and destroy the postsynaptic structures or disturb the Agrin/lipoprotein receptor-related protein 4 (LRP4)/muscle-specific tyrosine kinase (MuSK)/Dok7/rapsyn pathway. These processes lead to an impaired neuromuscular junction transduction characterized by fluctuating muscle weakness and fatigability (5,6). The muscle weakness can be generalized or localized. The respiratory muscles are less affected; however, their involvement can be life-threatening.

The following antibodies have been reported in MG: anti-acetylcholine receptor (anti-AChR), anti-muscle-specific tyrosine kinase (anti-MuSK), anti-lipoprotein receptor-related protein 4 (anti-LRP4), anti-agrin, anti-titin, anti-ryanodine receptor (anti-RyR), anti-collagen Q, and anti-voltage-gated potassium channel (anti-KV1.4) antibodies (Figure 1). MG is a heterogeneous disease in terms of pathophysiology, depending on the antibody status and other factors (Table 1). Serological tests are important in the management of patients with this disease because of their potential implications: (I) autoantibodies against the postsynaptic membrane are the most important biomarkers for the diagnosis of MG as they are widely available and antibody detection is important for distinguishing congenital myasthenia from MG; (II) patients with MG can be stratified into several subgroups: anti-AChR-positive MG (including early-onset and late-onset MG, and thymoma-associated MG), anti-MuSK-positive MG, anti-LRP4-positive MG, and mixed antibodies-positive MG. 

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onset), anti-MuSK-positive MG, anti-LRP4-positive MG, seronegative MG, thymoma-associated MG, and ocular MG. The antibody status is an important basis for MG classification; (III) the treatment regimen for MG varies according to the different antibodies. For example, patients with an anti-MuSK-positive MG have a poor response to oral acetylcholinesterase inhibitors. In fact, these medications may even be deleterious in these patients. 3,4-diaminopyridine is suggested as a potential substitute in these cases (7–9). Plasma exchange is usually the preferred over intravenous immunoglobulin treatment in patients with deteriorating anti-MuSK-positive MG; (IV) because the level of antibodies decreased in most patients with MG after plasma exchange or immunosuppressants therapy, the antibody levels might be useful as a monitoring indicator for therapy (10–12), although the antibody level has a limited value in monitoring of the disease due to the heterogeneity of the antibodies; (V) the antibody status is one of the factors influencing the outcome in patients with MG (13). Patients with ocular MG with anti-AChR or anti-MuSK antibodies were prone to develop generalized MG (14,15), whereas double-negative patients, those without anti-AChR or anti-MuSK antibodies, had a better prognosis (13). In summary, serological testing is crucial for diagnosis confirmation and therapy decision in patients with MG and is therefore recommended in the management of these patients.

**Anti-AChR antibodies**

Anti-AChR antibodies are the most common and highly specific antibodies in MG, found in 70–85% of all patients with MG (5,6,16,17), in 85–90% of patients with generalized MG, and in only 50–60% of patients with ocular MG (18–22). In the Chinese population, anti-AChR antibodies were present in 50–91.2% of patients with
ocular MG, in 82.4–86.4% of patients with generalized MG, and in 72–86.2% of all patients with MG (23-26). These results are similar to the reported prevalence of anti-AChR antibodies worldwide, even though one group reported a higher prevalence (91.2%) of these antibodies in patients with ocular MG (23). This variation may be due to the difference in the assays and the enrolled population. The pathogenicity of the anti-AChR antibodies has been well understood. These antibodies disturb the transmission in the neuromuscular junction primarily by complement binding and activation, leading to damage of the post synaptic membrane, by crosslinking to the AChRs, leading to accelerated endocytosis and degradation of these receptors, and direct blockade of the AChR binding sites also involved (27-31). Anti-AChR antibodies mostly belong to the IgG1 and IgG3 subtypes and primarily target the main immunogenic region (32). The antibodies against the α subunit of the AChRs are more pathogenic than those against the β subunit (33). In most studies, there was no correlation between the anti-AChR antibody concentration and the disease severity among different individuals (5,6,16,17). However, these studies focused on the total anti-AChR antibody level. Considering that the epitope variability influences the disease severity (33), identifying the binding region and subclass of antibodies may improve the correlation. The levels of antibodies against the main immunogenic region had a stronger correlation with the disease severity than those of the total anti-AChR antibodies (34). The value of repeated anti-AChR antibody testing in patients with MG is controversial. It may be useful for evaluation of the short-term therapy effect, particularly after plasma exchange. A correlation has also been noted between the changes in the anti-AChR antibody concentration and the disease severity. A strong correlation existed between the change in anti-AChR antibody concentration and the change in the clinical condition during immunosuppressive treatment and after thymectomy, whereas only a weaker correlation was present in periods without immunosuppression (35). In another report, there was a longitudinal association between the concentration of anti-AChR antibodies and the Myasthenia Gravis

Table 1 Clinical characters in different antibodies status

<table>
<thead>
<tr>
<th>Antibodies status</th>
<th>Prevalence</th>
<th>Clinical characters</th>
<th>Electrophysiology</th>
</tr>
</thead>
<tbody>
<tr>
<td>AchR-Ab</td>
<td>70–85%</td>
<td>Most typical</td>
<td>Positive most frequently</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Weakness fluctuate over time and induced by exercise</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Response well to AChEIs</td>
<td></td>
</tr>
<tr>
<td>MuSK-Ab</td>
<td>1–10%, may variation in different areas</td>
<td>Female predominant</td>
<td>Low yield of RNS in conventional limb muscles, but lager percentage of in facial nerve</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Tend to have cranial and bulbar involvement, limb weakness is less common</td>
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<tr>
<td></td>
<td></td>
<td>Fluctuation of weakness is less</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>Poor response to AChEIs</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Prone to be refractory MG</td>
<td></td>
</tr>
<tr>
<td>LRP4-Ab</td>
<td>1–5%, may variety depend on ethnic</td>
<td>Isolated ocular MG or mild generalized symptoms</td>
<td>RNS is infrequently abnormal, and jitter values were lower</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Rarely develop a crisis</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Response well to AChEIs</td>
<td></td>
</tr>
<tr>
<td>Titin-Ab</td>
<td>27–28.4% in all MG and 95% in thymomatous MG</td>
<td>Indication of thymoma</td>
<td>No report</td>
</tr>
<tr>
<td>RyR-Ab</td>
<td>14–23.8% in all MG and 70% in thymomatous MG</td>
<td>Indication of thymoma</td>
<td>No report</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Indication of severe MG</td>
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</table>

Ab, antibody; AChR, acetylcholine receptor; MuSK, muscle-specific kinase; LRP4, lipoprotein receptor-related protein 4; RyR, ryanodine receptor; MG, myasthenia gravis; AChEIs, acetylcholinesterase inhibitions; RNS, repetitive nerve stimulation.
Foundation of America (MGFA) classification in the subgroup with immunosuppressive treatment (36). These two studies adopted the MGFA classification to evaluate disease severity. Recently, the MG severity score was also used for assessment of the clinical condition. No significant correlation was demonstrated between the absolute change in the anti-AChR antibody titers and any severity measure; however, there was a strong correlation between the percentage change in the antibody level and the change in the manual muscle testing grade (12,37).

Radioimmunoprecipitation assay (RIPA)

RIPA is the most rigorously validated test for anti-AChR antibody detection. It was first applied to detect anti-AChR antibodies in 1976 (20). First, the AChRs are labeled with $^{125}$I-$\alpha$-bungarotoxin, which has a high affinity for the nicotinic AChRs. After incubation with the serum to be tested, a second antibody is added to precipitate the $^{125}$I-$\alpha$-bungarotoxin-anti-AChR complex. Washing steps should be performed to prevent a high background after the AChR labelling and incubation steps. The precipitate is counted and the precipitation rate is compared to that of healthy control samples (Figure 2). The AChRs were initially obtained from human amputated limb muscle until Sine et al. developed a new way to produce functional human AChRs from the TE671 cell line in 1988 (38). However, the AChRs derived from TE671 cells cannot replace the human muscle AChRs because the overall sensitivity was 7% less when using TE671 cell-derived AChRs (39). This is due to the fact that there are two subtypes of human muscle AChRs: fetal ($2\alpha\beta\gamma\delta$) and adult ($2\alpha\beta\varepsilon\delta$). The TE671 cells mainly express the fetal subtype. However, in some patients with ocular symptoms or low anti-AChR antibody titers, the antibodies mainly target the adult AChR subtype. To solve this problem, further improvement was made in 1996 by stably transfecting TE671 cells with cDNA encoding $\varepsilon$ subunit (TE671-$\varepsilon$) to produce a high concentration of adult AChRs (40,41). At present, RIPA for serum anti-AChR antibodies is mainly based on $^{125}$I-$\alpha$-bungarotoxin-labelled AChRs, which are mixed, containing both fetal and adult form subtypes. A carefully balanced mixture of detergent-solubilized fetal and adult forms of the receptor is the optimum. Currently, the overall diagnostic sensitivity of the routine RIPA to detect anti-AChR antibodies is 82% (42), and its specificity is very high, up to 98%. Because of its high sensitivity and specificity as well as widespread use, RIPA is considered as the "golden standard" for anti-AChR antibody testing. In addition, titres can be accurately determined with this method and
the assay can be performed quickly, in 1-2 days. There are commercially available testing kits, such as RiaRSR™ AChRAb from RSR limited (United Kingdom), ARAb RIA from IBL international (Germany), AChRAb RIA from Diasource (Belgium). The main disadvantage of RIPA is its radioactivity.

Enzyme-linked immunosorbent assay (ELISA)

ELISA for anti-AChR antibodies was first performed by Furukawa et al. (43). The $^{125}$I-α-bungarotoxin was replaced by HRP-linked α-bungarotoxin and the results were similar to those obtained using RIPA. However, this method was not widely adopted. Hewer et al. developed the competitive ELISA (44). The mAb1 antibodies, a type of monoclonal antibodies against AChRs (mixture of fetal and adult), were used to coat the ELISA plate wells, and the other two types, mAb2 and mAb3, were labeled with biotin. In the presence of anti-AChR antibodies, a sandwich is formed between the coated MAb1 antibodies, the anti-AChR antibodies, and the biotin-labeled mAb2 and mAb3 antibodies. When anti-AChR antibodies from the test serum are included, the formation of this sandwich is inhibited and the amount of biotinylated mAb antibodies bound to the plate wells (via AChRs and immobilized mAb antibodies) is reduced. The higher the concentration of anti-AChR antibodies in the test serum, the greater the inhibition of the mAb-biotin binding [detected by the addition of streptavidin-peroxidase (SA-POD)], thus providing a quantitative anti-AChR antibody ELISA (Figure 2). There was a good agreement between RIPA and ELISA for the anti-AChR antibody measurement ($r=0.85; n=83; P<0.001$). This method has a sensitivity and a specificity at least as high as those of RIPA. ELISA is a non-radioactive assay, and as it has a similar performance to that of RIPA, it can be a good alternative for laboratories where radioactive assays cannot be performed. There are commercially available inhibition ELISA kits, such as that by RSR (United Kingdom).

Cell-based assay (CBA)

The CBA for anti-AChR antibodies was developed in 2008 (45,46). Cultured human embryonic kidney (HEK) cells were transfected with the AChR subunit and rapsyn cDNAs. Rapsyn, which is a key component of the subsynaptic cytoskeletal complex, was added in order to obtain dense clusters of AChRs on the membrane of HEK cells. When a test serum is added, the antibodies bind to the clustered AChRs on the cell membrane. Subsequently, second, fluorescence-labeled antibodies are added to detect the cell membrane-bound anti-AChR antibodies and the results can be analyzed by flow cytometry (Figure 2). CBA has a higher sensitivity than RIPA and a specificity of nearly 100% (47). It detected low-affinity anti-AChR antibodies in 38.1–66% of patients with MG who were previously regarded as being negative on RIPA (45-47). In addition, 50% of the previously seronegative (anti-AChR-negative) patients with ocular MG were found to be anti-AChR-positive when tested with CBA (45). The low-affinity anti-AChR antibodies were all of the IgG1 subtype and pathogenic in an animal model, indicating their similar mechanisms. CBA is a non-radioactive assay with a higher sensitivity and specificity. Furthermore, the antigens are expressed in a clustering form on a cell surface, which is more similar to the antigens in vivo; hence, it is more a sensitive method than the RIPA and ELISA assays with purified AChR molecules. CBA can offer a semi-quantitative estimation of autoantibody titers and quantification can be easily performed with flow cytometry. However, it is more laborious and there is no commercial kit available at present.

Other assays

Fluorescence immunoprecipitation assays (IFPA) was developed in Oxford, UK, and Yang et al. applied it in Chinese patients for anti-AChR antibody detection (48). Transfected HEK 293 cells were used to produce fetal and adult AChRs tagged with fluorescent proteins [enhanced green fluorescence protein (EGFP)]. The AChR-EGFP are mixed with serum from patients and then protein A sepharose beads are added to catch the antibody-receptor complexes. After rotation and washing, the beads are transferred to conical bottom plates and the fluorescence units values are read using a fluorescent plate reader to estimate the titers of anti-AChR antibodies. IFPA has a good agreement with RIPA ($r^2=0.6; P<0.0001$). Although, IFPA was found not to be as sensitive as RIPA (76% vs. 80%, respectively), it may be due to the relatively small sample size. The advantage of this approach is that it can use different fluorescence-tagged proteins (such as mixed with MuSK-mcherry) for detection of antibodies against different antigens in the same sample.
Anti-MuSK antibodies

Anti-MuSK antibodies were first described in anti-AChR-negative patients with MG in 2001 (49,50). They can be detected in 1–10% of all patients with MG and in 13–41% of anti-AChR-negative patients (51,52). This variety depends on the geographical location (53-55). Among the Chinese population, it was found in 4% of anti-AChR-negative patients with MG and in 2.3–5.9% of all patients with MG (24,25,48). Anti-MuSK antibodies are predominantly IgG4 antibodies that do not bind complement. Hence, they were once considered to be “benign antibodies” or bystanders. Klooster et al. proved the pathogenicity of these antibodies in mice and observed a reduced density and a fragmented area in the neuromuscular junction AChRs (56). Huijbers et al. and Koneczny et al. elucidated the underlying pathogenic mechanism. Anti-MuSK-IgG target a structural epitope in the first Ig-like domain of MuSK, preventing the binding between MuSK and Lrp4, thereby inhibiting the agrin-induced clustering (57,58). In summary, anti-MuSK antibodies disturb the AChR cluster formation and aggregation on the postsynaptic membrane rather than decreasing or blocking AChRs (16,59). In contrast to the anti-AChR antibodies, both in individual cases and in the whole population, a correlation was found between the levels of antibodies and the disease severity (60). The correlation may be because, unlike heterogeneity of AChR-Abs, the majority of MuSK-Abs targets the first Ig like domain. Patients with anti-MuSK antibodies were prone to have refractory MG and were more likely to develop respiratory weakness (50,61). Their response to acetylcholinesterase inhibitors is often insufficient or their condition may even exacerbate after administration of pyridostigmine. Furthermore, these patients respond better to plasma exchange than to intravenous immunoglobulin treatment (61).

RIPA

RIPA is regarded as a standard test for anti-MuSK antibodies as well. The procedure is similar to that of RIPA for anti-AChR antibodies. Antigens labeled with $^{125}$I were derived from engineered HEK 293 cells expressing human MuSK ectodomain or recombinant rat MuSK (51,62). The $^{125}$I-MuSK is used to precipitate the anti-MuSK antibodies. Using this method, 13–41% of anti-AChR-seronegative patients were found to be anti-MuSK-positive. Its specificity is speculated to range from 97% to 100%. Commercial RIPA kits are available, including that by RSR (United Kingdom) and CISbio (France). Moreover, Trakas et al. developed a two-step RIPA, using immobilized antigens to enrich the antibodies before the routine RIPA (63). This two-step RIPA has a higher sensitivity, but there is no available commercial kit.

ELISA

ELISA was the first reported anti-MuSK-antibody assay (49). Antigens were derived from MuSK-transfected COS7 cells and were coated on the plate. Plasmas from patients are added. The bound antibodies are detected by horseradish peroxidase (HRP), followed by substrate and measuring by absorbance. ELISA is a substitute for RIPA for anti-MuSK antibodies detection (64). A commercial ELISA kit can be acquired from IBL international (German). Their measurements of MuSK autoantibodies showed that in 253 serum samples sensitivity of ELISA is up to 95.8% and specificity is 100% compare to RIPA (https://www.ibl-international.com/en/musk-ab-elisa).

CBA

CBA was also applied in the diagnosis of MG with anti-MuSK antibodies. HEK 293 cells transfected with full-length human MuSK cDNA were used for anti-MuSK antibody detection (46,65). CBA is more sensitive than RIPA or ELISA, and its specificity is estimated to be 97–98% (46,65,66). Furthermore, the CBA-detected anti-MuSK-IgGs were capable of disrupting the agrin-induced AChR clustering in myotube cultures (46,65). No commercial kit is available at present.

Other assays

IFPA was used for anti-MuSK antibody measurement in China by Yang et al. (24,25,48). The extracellular domain of human MuSK, covalently tagged with mcherry, was expressed and secreted by stably transfected insect cell line S2 and then purified through affinity column. The purified MusK-mcherry were mixed with AChR-EGFP and used to measure both antibodies in the same assay. For anti-MuSK antibody detection, IFPA seems to be in agreement with RIPA, but less sensitive than CBA (two
patients were positive for anti-MuSK antibodies by both RIPA and FIPA, but CBA was identifying in additional three patients).

**Anti-LRP4 antibodies**

Anti-LRP4 antibodies were first found in patients with MG in 2011 (67-69). They can be found in 1–5% of all patients with MG or 7–32.7% of the “serum double negative” patients (70). The variation may be due to ethnic differences (70). Two groups have investigated anti-LRP4 antibodies in the Chinese population. Li et al. reported that 2 of all 116 (1.7%) patients with MG and 2 of 50 (4%) double-seronegative patients were positive for anti-LRP4 antibodies (71). Hong et al. found that 7% of the double-seronegative patients with MG were anti-LRP4-positive (25). The prevalence of anti-LRP4 antibodies in China is similar to that Japan, but less than that in Poland (67,70). These antibodies can co-exist with anti-AChR or anti-MuSK antibodies and can be detected in other neuroimmune diseases (neuromyelitis optica and multiple sclerosis) (69,70). Anti-LRP4 antibodies were also found in 14.9–23.1% of patients with amyotrophic lateral sclerosis (ALS). One possible explanation is that the antibodies are generated owing to immune dysfunction in some patients with ALS (72). LRP4 forms a multiprotein complex with MuSK, participating synapse formation and AChR clustering via the agrin/LRP4/MuSK/Dok7/rapsyn pathway (16,73). Anti-LRP4 antibodies block the agrin–LRP4 interaction, disturbing the AChR clustering in the membrane. The pathogenicity of anti-LRP4 antibodies has been proved in an animal model (74). Anti-LRP4 antibodies are predominantly of the IgG1 and IgG2 subtypes (70,75). In contrast to patients with anti-MuSK antibodies, most of these patients present with isolated ocular or only mild generalized MG and rarely experience crisis (67,68,76). Thus, in most reports anti-LRP4 antibodies seem to be indicating a more favorable prognosis. However, Higuchi et al. found that a severe limb muscle weakness or a progressive bulbar palsy or both appeared in anti-LRP4-positive patients with MG (67). The fact that some of those patients were anti-MuSK-positive may partly explain this difference.

**ELISA**

Zhang et al. reported anti-LRP4 antibodies detection through ELISA (69). Briefly, immuno-plates were coated with purified recombinant LRP4, which was derived from HEK293 cells transfected with plasmid encoding full-length ecto-LRP4. The serum to be tested is added, followed by alkaline phosphatase–goat anti-human Ig secondary antibody. The activity of the immobilized alkaline phosphatase is measured to estimate the level of anti-LRP4 antibodies. This method found that 9.2% of double-seronegative patients with MG were anti-LRP4-positive. Its specificity is estimated to 98%. No commercial ELISA kit is available.

**CBA**

CBA was developed for anti-LRP4 antibodies to analyze the epidemiology and clinical characteristics of anti-LRP4-positive MG (70). HEK293 cells transfected with LRP4-GFP-loaded plasmid are used to bind the anti-LRP4 antibodies. Subsequently, fluorochrome-conjugated anti-human IgG antibodies are added to estimate the level of anti-LRP4 antibodies semi-quantitatively. CBA is more sensitive than ELISA, and its specificity is near 98%. No commercial tests for anti-LRP4 antibodies are available.

**Anti-titin antibodies and anti-RyR antibodies**

Anti-titin and anti-RyR are antibodies against components of the skeletal muscular cells. As their antigens are intracellular molecules, it is controversial whether they have pathogenic significance in vivo. There was evidence that they can penetrate the cell membrane and bind to their target (77). They might be disease markers rather than mediators of muscle weakness (78) and present in 27–28.4% and 14–23.8% MG patients (24,79). Presently, anti-titin-antibody RIPA is a useful tool for serological diagnosis of serum triple-negative patients with MG (namely anti-AChR, anti-MuSK, and anti-LRP4 antibodies are all negative) (80). Of all triple-negative patients, 13.4% were also anti-titin antibody positive, 3.6% of patients with other neurological disease were positive, and none of the healthy controls or patients with myopathy. In addition, anti-titin and anti-RyR antibodies have a close relationship with thymoma; anti-titin antibodies appeared in 95% of patients with MG with thymoma and anti-RyR antibodies appeared in 70% of patients with MG with thymoma (79,81). The combination of anti-titin and anti-RyR antibody testing has a 95%
sensitivity and a 70% positive predictive value (79). The presence of these antibodies also indicates severe MG (79,82). In a few patients with MG, autoimmune-mediated myopathies developed, particularly in thymoma-associated MG. These patients with myopathies have autoantibodies against the molecules on skeletal muscles, including Titin-Ab and RyR-Ab. The mechanisms for the development of MG myopathy remain unclear, but it may predict a poor outcome (83,84).

Anti-titin and anti-RyR antibodies were detected by radioimmunoassay or ELISA or RIPA (24,25,85,86). In the Chinese population, these antibodies were found in 64% and 55% patients with MG, respectively, and they were present with a high frequency in MG with thymoma (24-26).

Other antibodies

Anti-agrin antibodies were reported in 2014 (87). Agrin regulates the formation and maintenance of the neuromuscular junction via the agrin/LRP4/MuSK/rapsyn pathway (16,73,88). Anti-agrin antibodies interfere with the function of agrin, leading to a neuromuscular junction impairment. Whether anti-agrin antibodies contribute to the muscle weakness is unclear. They can be detected only in a minority of patients with MG and are accompanied with other antibodies (such as AChR-Ab, MuSK-Ab) (87,89). Antibodies against collagen Q were detected in 3% patients with MG but have also been found in healthy controls (52,90). Their diagnostic or pathogenic value remains to be proven. Antibodies against KV1.4 channel are found in 10–20% of patients with MG. MG with KV1.4-Ab seemed to be of greater severity or with heart impairment (91,92). Thymoma diagnosed by imaging tests was more frequent in the anti-KV1.4-Ab positive patients than in the negative patients and expression of KV1.4 mRNA was found in the abnormal thymus tissue from patients with MG (93).

Clinical application of serological assay

The performance of serological tests in screened populations and the selection of reasonable methods are important for an accurate diagnosis. We would like to propose the following. First, serological tests should be performed in patients with clinically suspected MG. It means that before the serological test, a clinical evaluation should be performed. Screening of large unselected populations may decrease the positive predictive value of the diagnostic tests. Furthermore, a minority of patients with MG are still seronegative. Their diagnosis could not depend solely on a serological test. In addition, some of these markers can be found in other diseases (such as LRP4-Ab in neuromyelitis optica, multiple sclerosis and ALS). Thus, it is the combination of a positive serological test (including AChR-Ab, MuSK-Ab, LRP4-Ab) and highly suspicious clinical manifestations that establish the MG diagnosis. Second, the proper markers should be selected according to the practical condition. Nine antibodies associated with MG have been reported and new antibodies may be discovered in the future. The detection of all kinds of antibodies for every patient with MG is expensive and unnecessary. Anti-AChR, anti-MuSK, and anti-LRP4 antibodies are the most important markers since they are the most common pathogenic antibodies and are not only an important clue for diagnosis but are also the basis for clinical classification and further guide therapy decisions. Not every patient needs to undergo all three antibody tests, because MuSK-Ab and AChR-Ab rarely appear in the same patient. We recommend that suspected patients undergo AChR-Ab detection as a first serologic test. MuSK-Ab detection should be performed when AChR-Ab is negative or the patient is highly suspected to have MuSK-associated MG. If the patient is double-negative, the clinical assessment should be reviewed. One should also consider to test other antibodies such as LRP4-Ab, and perhaps also Titin-Ab and Agrin-Ab if. The procedures are displayed in Figure 3. RIPA, ELISA, CBA are popular methods for serological diagnosis of MG and each method has distinct characteristics (Table 2). The RIPA is still the routine assay, with advantages on specificity, sensitivity and reliability. ELISA could be a good alternative to RIPA for institutions that do not perform RIPA. CBA is more sensitive but it is laborious and is not widely adopted. CBA is suitable to detect antibodies in the samples from patients highly suspected by the neurologist but RIPA are negative. CBA also can be used for detection of LRP4.

Conclusions

Antibody tests are even more important than before in the diagnosis of MG because of the development of assays with a higher sensitivity and the identification of novel antigens in MG. Furthermore, treatment decisions for MG should
Suspected MG

AChR-Ab test, LRP4-Ab test

Patients with MG

Titin-Ab, RyR-Ab test (optional)
CT scan of thymus

Patients with MG

MuSK-Ab test

Not MG

Clinical assessment:
Clinical features
Rest test, ice test
Neostigmine test

Suspected MG

Not MG

CT scan of thymus

Patients with MG

MuSK-Ab test

Not MG

MG

Not MG

Other antibodies test (optional):
Titin-Ab, RyR-Ab, Agrin-Ab, etc

Patients with MG

positive

negative

positive

negative

MuSK-Ab test

Figure 3 The procedures for detecting MG associated antibodies. Suspicious patients accepting AChR-Ab and LRP4-Ab detection at first serologic test as AChR-Ab is the most common type while LRP4-Ab may co-exist with AChR-Ab or MuSK-Ab. MuSK-Ab detection should be performed when AChR-Ab is negative or the patient is highly suspected to be MuSK-associated MG. If the patient is triple-negative, the clinical assessment should be reviewed and consider to test other antibodies. MG, myasthenia gravis; Ab, antibody; MuSK, muscle-specific kinase; AChR, acetylcholine receptor; LRP4, lipoprotein receptor-related protein 4; RyR, ryanodine receptor; RNS, repetitive nerve stimulation; SFEMG, single-fiber electromyography.
be personalized; accurate MG subgrouping based on the antibody status is an important step.

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Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

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RIPA, radioimmunoprecipitation assay; ELISA, enzyme-linked immunosorbent assay; CBA, cell-based assay; MG, myasthenia gravis.

<table>
<thead>
<tr>
<th>Table 2 Methods for measuring MG associated antibodies</th>
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<td><strong>RIPA</strong></td>
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<td><strong>Advantages</strong></td>
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<td><strong>Disadvantages</strong></td>
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41. Beeson D, Jacobson L, Newsom-Davis J, et al. A transfected human muscle cell line expressing the adult subtype of the human muscle acetylcholine receptor