Myasthenia gravis during pregnancy: what care should be taken?

Miastenia gravis durante a gestação: quais cuidados devem ser realizados?

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ABSTRACT
Myasthenia gravis (MG) is an autoimmune disease in which the peak incidence is among women of childbearing age. For this reason, there is an overlap between the occurrence of this disease and pregnancy. It is known that MG symptoms can worsen during pregnancy and postpartum, and that pregnancy has special characteristics in MG patients. Children born to myasthenic mothers are at risk of having transient neonatal myasthenia. We briefly review the main relationships between MG and pregnancy, and we make recommendations for MG therapy, pregnancy, delivery, breastfeeding and newborns.

Keywords: Myasthenia gravis; Pregnancy; Delivery; Breastfeeding; Transient Neonatal Myasthenia; Therapy.

RESUMO
Miastenia gravis (MG) é uma doença autoimune cujo pico de incidência nas mulheres coincide com a idade fértil. Por esse motivo, existe uma sobreposição entre a ocorrência desta doença com a gestação. Sabe-se que os sintomas da MG podem piorar durante a gestação e no puerpério, e que a gestação possui características especiais em pacientes com MG. Existe também um risco de miastenia neonatal transitória em recém-nascidos de gestantes miastênicas. Nós revisamos brevemente as principais relações entre MG e gestação, e realizamos recomendações para o tratamento da MG durante a gestação, os cuidados durante a gravidez, incluindo a via de parto, e com o recém-nascido e com a amamentação.

Palavras-chaves: Miastenia gravis; Gestação; Via de parto; Amamentação; Miastenia Neonatal Transitória; Tratamento.

INTRODUCTION
Myasthenia gravis (MG) is an autoimmune disorder that is caused by antibodies against the neuromuscular junction. Its main characteristic is extraocular and proximal muscle weakness associate with fatigability¹,². The peak incidence of MG in women coincides with childbearing age³–⁵. Prenatal counseling should be provided to all MG patients who are women in the reproductive age group. MG does not affect patients' fertility⁴, but it does interfere with family planning⁴. Many patients have abstained from having children mainly because of fear of the possible effects of MG medications on the development of the unborn child⁴. Thus, questions have arisen concerning the impact of MG on pregnancy, as well as of pregnancy on MG; and concerning the effect of medications used to treat MG on fetal development, mode of delivery, safety of breastfeeding and newborn care.

In this brief review, we describe the main relationship between MG and pregnancy and we make recommendations for MG therapy, pregnancy, delivery, breastfeeding and newborns.

THE EFFECT OF PREGNANCY ON MYASTHENIA GRAVIS
In 12% to 15% of women with MG, the first manifestations of the disease were during pregnancy or in the postpartum
A population-based case-control study indicated that women in the postpartum period were at higher risk of experiencing symptoms of MG, especially after the first pregnancy. If a diagnosis of MG is suspected during pregnancy or the postpartum period, an investigation should be carried out in a similar way to that performed outside these periods (MG has the same diagnostic criteria during pregnancy as outside it). The main MG diagnostic tools are detection of serum antibodies (against acetylcholine receptor [AchR] and muscle-specific tyrosine kinase [MuSK], among others) and abnormalities in electrophysiological tests (repetitive nerve stimulation and single fiber electromyography).

The view that pregnancy does not influence the evolution of MG over the long term seems to be well-established in the international literature. With regard to the effects of pregnancy on MG over the short term, the symptoms may improve, remain stable or worsen, and these effects can vary from pregnancy to pregnancy in the same patient. In a Brazilian study, the condition of 50% of the patients worsened during pregnancy, mainly in the second trimester, and in the postpartum. Other studies have suggested that the first trimester is also a critical period. It is believed that the third trimester is a period of greater stability or even clinical improvement for these patients. This is perhaps the result of increased production of α-fetoprotein, thus inducing physiological immunosuppression, since α-fetoprotein is believed to be an effective inhibitor of AchR antibodies. In contrast, the postpartum period would be the most critical because of the production of cytokines and immunoglobulins induced by estrogen and decreased levels of α-fetoprotein, the risk of infection and the fatigue induced by stress and sleep deprivation related to the new family routine. MG symptom exacerbations may occur even among patients with good control prior to conception. In a recent study, a myasthenic crisis during pregnancy or the postpartum period would be the most critical because of the production of antibodies against MuSK has greater impact on pregnancy or greater risk of worsened myasthenic symptoms than does presence of antibodies against AchR. Retrospective studies have also demonstrated that prior thymectomy had no effect on myasthenia symptoms during pregnancy.

Several studies have shown a higher rate of preterm premature rupture of amniotic membranes (PPROM) and prematurity, possibly associated with use of corticosteroids. The abortion rate is similar to that of the general population and is not influenced by MG treatment. Among the possible complications of pregnancy, preeclampsia occurs in 6–8% of all pregnancies. An association between preeclampsia and MG is rare, but management of such cases is challenging. Hypertension should be treated with methyldopa and oral hydralazine. If systolic pressure is greater than 160 mmHg or diastolic pressure is greater than 110 mmHg, intravenous hydralazine is indicated. Beta blockers and calcium channel blockers should be avoided due to the potential risk of worsening MG symptoms. Magnesium sulphate is the gold standard therapy for eclampsia and severe preeclampsia; however, it should not be used in myasthenic patients because it inhibits presynaptic calcium influx at the neuromuscular junction. Options for preventing seizures include levetiracetam and valproic acid, but the latter should be avoided in cases of major hepatic impairment. For refractory cases, use of phenytoin may be indicated, although there is a theoretical risk of exacerbating myasthenic weakness by reducing the sensitivity of acetylcholine receptors. Benzodiazepines can be used for seizure control with caution, due to muscle relaxant effects via central potentiation of gamma amino butyric acid (GABA) release.

The delivery route in these patients needs to be an obstetric indication. In other words, the diagnosis of MG does not contraindicate normal delivery per se. This is because the myometrial muscle is composed of smooth muscle and is not affected by MG autoantibodies. Patients may worsen in the second stage of labor when the striated muscles are involved and, at that time, maternal exhaustion may require use of forceps or vacuum extraction. Studies have shown cesarean rates ranging from 11% to 78.3%. An association between cesarean sections and MG is rare, but management of such cases is challenging. Hypertension should be treated with methyldopa and oral hydralazine. If systolic pressure is greater than 160 mmHg or diastolic pressure is greater than 110 mmHg, intravenous hydralazine is indicated. Beta blockers and calcium channel blockers should be avoided due to the potential risk of worsening MG symptoms. Magnesium sulphate is the gold standard therapy for eclampsia and severe preeclampsia; however, it should not be used in myasthenic patients because it inhibits presynaptic calcium influx at the neuromuscular junction.

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use of non-opioid drugs such as acetaminophen and nonsteroidal anti-inflammatory. Use of opioid/narcotic analgesics such as tramadol or morphine should be avoided due to the possible risk of respiratory depression.

**THERAPEUTIC POSSIBILITIES FOR MYASTHENIA GRAVIS DURING PREGNANCY**

Ideally, treatment for pregnant myasthenic patients should consist of use of corticosteroids and pyridostigmine. If possible, pregnancy planning should be implemented, with enough time to assess the risks and benefits of each medication and to make therapeutic changes. New drugs should preferably not be started during pregnancy. Breastfeeding, on the other hand, should be stimulated, although it should be avoided at the peak of the medication dose.

Pyridostigmine is an anticholinesterase agent that is used for symptomatic management of MG. There are no reports of fetal malformations, and breastfeeding is safe.

Corticosteroids have been used for many years in pregnant women with other diseases. Their use has been correlated with increased risk of cleft palate (in a single study in which they were applied in the first trimester of pregnancy). Adrenal suppression in newborns has been reported. Corticosteroids are safe during breastfeeding, and there are no reports of adrenal suppression in newborns.

Several studies on pregnant women with MG using azathioprine have already been carried out. For some authors, azathioprine is associated with prematurity, intrauterine growth retardation and higher risk of teratogenesis (prevalence between 0% and 11.8%). However, others have reported that azathioprine is safe during pregnancy and during breastfeeding, and they have stated that it should be the immunosuppressant drug of choice, after corticosteroids, in cases of MG during pregnancy. Azathioprine is considered to be a safe drug for use during pregnancy in Europe, whereas it is still contraindicated in the United States and in Brazil. It is known that the fetal liver does not produce pyrophosphorylase, the enzyme responsible for converting azathioprine into its active metabolites.

Use of cyclosporine is associated with increased risk of gestational diabetes, gestational hypertensive disease, prematurity and low birth weight. However, this drug is considered safe during breastfeeding, according to some authors.

Use of tacrolimus is also associated with a higher risk of prematurity and low birth weight, with divergent results in relation to teratogenesis. An in vitro study suggested that there was higher risk of teratogenesis if tacrolimus was used in association with prednisolone. Some authors have stated that breastfeeding should be avoided by patients who are using this medication.

Other immunosuppressive drugs should be discontinued during pregnancy and the breastfeeding period (e.g., mycophenolate mofetil, methotrexate and cyclophosphamide). Both mycophenolate mofetil and methotrexate must be discontinued due to the increased risk of miscarriage and teratogenesis. Mycophenolate mofetil should be suspended and replaced ideally six weeks before the patient becomes pregnant, while methotrexate should be suspended and replaced three months before pregnancy. Breastfeeding is not recommended while using either of these medications. Similarly, cyclophosphamide must be avoided during pregnancy due to the risk of teratogenesis; breastfeeding is also contraindicated. Cyclophosphamide should be suspended and replaced one year before pregnancy.

There are few case reports regarding the use of rituximab in pregnant women. It has been suggested that there should be an interval of 12 months between the last dose of rituximab and pregnancy. Breastfeeding is not recommended because of a lack of data on the possible effects of this drug. Eculizumab should be avoided during pregnancy and breastfeeding for the same reason.

If myasthenic crisis occurs, both immunoglobulin and plasmapheresis appear to be safe. There are studies on myasthenic pregnant patients who used immunoglobulin without complications, including as maintenance monotherapy. With regard to therapeutic plasma exchange (TPE), studies have mainly been conducted on pregnant women with other diseases. However, a recent study on 57 pregnancies, among which three were myasthenic, showed that therapeutic apheresis was well tolerated. There is a tendency to prefer immunoglobulin in such cases due to the possibility of hypotension during the TPE. If TPE is chosen, some precautions should be taken, such as: positioning the patient in the left lateral position to avoid compression of the inferior vena cava; adjustment of plasma volume in the second and third trimesters of pregnancy to compensate for the increase in maternal blood volume; the volume of blood in extracorporeal circulation should not decrease the blood volume by more than 10%, in order to avoid hypotension; preventing and monitoring for hypocalcemia; monitoring the levels of fibrinogen, because a decrease carries a greater risk of bleeding (and for that reason a minimum 24-hour interval between the last plasmapheresis session and the cesarean delivery is suggested); and determination of Rh status and consideration of possible Rh Ig re-administration after TPE, if necessary.

Table 1 indicates the main drugs used in treating MG, in relation to their safety profile for pregnancy and breastfeeding. It is also important to advise MG patients on the medications that are contraindicated due to MG (Table 2).
Table 1. Risk categories of medications used to treat myasthenia gravis during pregnancy and breastfeeding.

<table>
<thead>
<tr>
<th>Drugs</th>
<th>FDA pregnancy category</th>
<th>Breastfeeding category</th>
</tr>
</thead>
<tbody>
<tr>
<td>Azathioprine</td>
<td>D</td>
<td>Conflicting results; to be used with caution, as specified by the Brazilian Ministry of Health</td>
</tr>
<tr>
<td>Cyclophosphamide</td>
<td>D</td>
<td>Not advised</td>
</tr>
<tr>
<td>Cyclosporine</td>
<td>C</td>
<td>Conflicting results; to be used with caution, as specified by the Brazilian Ministry of Health</td>
</tr>
<tr>
<td>Eculizumab</td>
<td>C</td>
<td>Not advised because of lack of information</td>
</tr>
<tr>
<td>Immunoglobulins</td>
<td>C</td>
<td>Allowed</td>
</tr>
<tr>
<td>Methotrexate</td>
<td>X</td>
<td>Not advised</td>
</tr>
<tr>
<td>Mycophenolate mofetil</td>
<td>D</td>
<td>Not advised</td>
</tr>
<tr>
<td>Pyridostigmine</td>
<td>C</td>
<td>Allowed</td>
</tr>
<tr>
<td>Rituximab</td>
<td>C</td>
<td>Not advised because of lack of information</td>
</tr>
<tr>
<td>Corticosteroids</td>
<td>C</td>
<td>Allowed</td>
</tr>
<tr>
<td>Tacrolimus</td>
<td>C</td>
<td>Conflicting results; to be used with caution, as specified by the Brazilian Ministry of Health</td>
</tr>
</tbody>
</table>

FDA: United States Food and Drug Administration; D: positive evidence of risk; C: risk cannot be ruled out; X: contraindicated in pregnancy.

Table 2. Medications contraindicated for patients with myasthenia gravis.

Anesthetic agents
- Neuromuscular blocking agents

Antibiotics
- Aminoglycosides, fluoroquinolones, macrolides, metronidazole, nitrofurantoin, sulfonamides, tetracyclines and vancomycin

Cardiovascular drugs
- β-blockers, calcium channel blockers, quinidine and procainamide

Other drugs
- Amphetamines
- α-interferon
- Antihistamines
- Benzodiazepines
- Botulinum toxin
- D-penicillamine
- Iodinated contrast agents
- Lithium
- Magnesium, including milk of magnesia and antacids containing magnesium
- MAO inhibitors
- Monoclonal antibodies such as nivolumab and pembrolizumab hydroxide and magnesium sulfate
- Morphine
- Neuroleptics
- Oxytocin
- Quinine

Other drugs have been reported to exacerbate MG. The risks and benefits of starting a new medication must be considered carefully in relation to these patients, who should be followed up closely.

RISKS FOR NEWBORNS FROM MYASTHENIC MOTHERS

With regard to newborns, the main complication is a higher risk of premature birth. In MG, because of the antibodies against AChR, transient neonatal MG (TNMG) occurs at rates ranging from 10 to 20%. This does not relate to either maternal MG status or maternal AChR antibody concentration. It means that asymptomatic pregnant women are also at risk of having newborns with TNMG and, on the other hand, that most pregnant women with severe MG will have newborns without TNMG. It is believed that the occurrence of TNMG depends not only on maternal antibody levels, but also on antibody transportation across the placenta, antibodies binding to the child’s postsynaptic muscle membrane and safety factors in the neuromuscular junction. In addition, there have also been reports of TNMG in newborns from MG mothers with antibodies against MUSK and lipoprotein receptor-related protein 4 (LRP4), and also in antibody-negative MG patients. Some studies have suggested that TNMG occurrence is lower when the pregnant woman has previously undergone thymectomy.

In most cases of TNMG, the newborn shows symptoms of hypotonia, ptosis, ophthalmoparesis and bulbar and respiratory weakness from within 12 hours after birth to several days later. TNMG develops as a result of maternal IgG antibodies that cross the placental barrier. The delay in onset
of symptoms is due to clearance of the anticholinesterase agents taken by the mother. The diagnosis is clinical and does not require measurement of autoantibodies or electrophysiological studies in the newborn\textsuperscript{15}. Symptoms can persist for periods from two weeks to months, and 90% recover within two months. Treatment consists of support in mild to moderate cases and immunoglobulin or plasmapheresis in severe cases. Cases of arthrogryposis and fetal AChR inactivation syndrome have also been reported, but both of these syndromes are rare. They are caused by the mother’s antibodies against fetal-type AChR\textsuperscript{3,10,31,45–47}.

It is important to emphasize that breastfeeding, provided that it is not contraindicated by any medications being taken by the puerperal woman, is not contraindicated, even in cases of TNMG, arthrogryposis or fetal AChR inactivation syndrome. This recommendation is based on the fact that the level of maternal IgG antibodies in breast milk comprises only 2% of the serum level, and that an even smaller proportion is absorbed by the child\textsuperscript{31,48}. Some studies have suggested that the breastfeeding schedule should not coincide with the peak serum concentration of the medications used by the mother\textsuperscript{4,49}.

**RECOMMENDATIONS**

The main recommendation is that prenatal counseling should be provided for all MG patients who are women in the reproductive age group. Preferably, these patients should use pyridostigmine and corticosteroids during pregnancy; use of mycophenolate, cyclophosphamide and methotrexate should be discontinued. Delivery needs to be carried out in accordance with obstetric rules. Anesthesia (epidural, spinal or local) is safe during delivery. Some drugs (e.g. neuromuscular blockers, oxytocin and magnesium sulfate) should not be administered to MG patients. The pediatric team needs to be prepared for the risk of transient neonatal MG. The family should be alerted to the possibility that MG could worsen in the postpartum period. Breastfeeding should be encouraged in most cases. Lastly, it should be noted that, despite all the above warnings, there is no increased risk of maternal or perinatal mortality among pregnant women with MG and, therefore, pregnancy plans should not, in most cases, be discouraged, provided that there is adequate monitoring of the patients.

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