

POriginal Research Article

Thymectomy in children with juvenile myasthenia gravis: Is it recommended?

Abstract:

Objective:

Juvenile myasthenia gravis (JMG) is a rare disease with possible severe forms. Thymectomy is supported by many authors particularly in generalised forms with positive AcetylCholine Receptor Antibody (AChR- Ab). The aim of our study was to discuss the indication and the outcome of thymectomy in five children with JMG.

Methods:

We performed a retrospective study of the medical files of patients with JMG, who were hospitalised in our pediatric surgery department for thymectomy. For all the children we performed Acetylcholine Receptor Autoantibody (AChR-Ab) test-system producer , a chest x-ray, a chest Computed Tomography Scan (CT-scan) and a Magnetic Resonance Imaging (MRI).

Thymectomy was indicated on the presence of anomalies on CT-Scan or MRI and realised via either thoracotomy or video-assisted thoracoscopy.

The study was conducted with the approval of the ethics committee at our institution.

Results:

Based on our findings, thymectomy in JMG with imaging anomalies seems to be associated with clinical improvement. In our patients complete remission after surgery was seen in one case, a clinical improvement with a decrease of the medical therapeutic dosage was noted in

three other cases. One patient could not be evaluated because of non adherence to medical treatment.

Surgery by thoracotomy or video assisted thoracoscopy (VAT) was also well tolerated by all the patients.

Conclusion:

The benefits of thymectomy in children with JMG is still controversial. However, many researchers have reported a sustained improvement of symptoms in the majority of patients after thymectomy. Our findings seem to encourage thymectomy in cases of JMG and particularly when imaging anomalies are found. VAT thymectomy was well tolerated and it is considered by many researchers to be the standard method for thymectomy in children.

Keywords: Juvenile myasthenia gravis, thymectomy, children, thoracoscopy.

Abbreviations and acronyms:

JMG: juvenile myasthenia gravis

AChR-Ab: acetylcholine receptor autoantibody

CT-Scan: computed tomography scan

MRI: magnetic resonance imaging

VAT: video assisted thoracoscopy

OMG: ocular myasthenia gravis

43 Introduction:

44 Juvenile myasthenia gravis (JMG) is a rare auto-immune disorder of neuromuscular
45 transmission caused by the production of antibodies against components of postsynaptic
46 membrane of the neuro-muscular junction (1). Young patients may present severe forms with
47 generalised muscle weakness with or without respiratory insufficiency. Specific treatment is
48 needed including immunosuppressive therapy and thymectomy particularly in presence of
49 thymus abnormalities (1-3). The aim of our study is to discuss indication and outcome of
50 thymectomy in children with JMG.

51 Patients and Methods:

52 We performed a retrospective study of the medical files of patients with JMG hospitalised in
53 our pediatric surgery department for thymectomy.

54 Diagnosis of JMG was established in pediatric departments based on a combination of clinical
55 symptoms and positive pyridostigmine test. Disease severity was evaluated according to
56 Osserman Classification. For all patients, we performed Acetylcholine Receptor Autoantibody
57 (AChRAb) test-system producer, a chest x-ray, a chest Computed Tomography Scan (CT-
58 scan) and a Magnetic Resonance Imaging (MRI).

59 All patients received a medical treatment which consisted of pyridostigmine and
60 corticotherapy.

61 Thymectomy was indicated in case of thymic anomalies on CT-Scan or MRI. It was realised
62 either via thoracotomy or with video-assisted thoracoscopy (VAT). VAT was introduced in
63 our department in 2009.

64

65 **Results:**

66 Five patients (table1) were included, 1 male and 4 females with sex ratio of 0.25. Mean age
67 was 7.7 years (2.5 to 14 years old). Four patients had a generalised JMG and one had an
68 ocular myasthenia gravis (OMG). Only one patient was AChR-Ab positive. Prior to
69 thymectomy, disease severity was graded as I for one patient, IIa for two patients and IIb for
70 two other patients.

71 Mediastinal enlargement was noticed on the chest x-ray of two patients (Fig1). The CT-scan
72 showed the presence of an eventual thymoma in three cases(Fig2) and a thymus hypertrophy
73 in the two other cases.

74 An **extended** thymectomy (Fig3) was indicated for all the patients. It was realised via antero-
75 lateral thoracotomy in three cases and via VAT in two cases.

76 The initial post operative course was complicated by a pneumothorax in case 2, related to a
77 pleural breach, successfully managed by assisted mechanical ventilation and pleural drainage.

78 Microscopic examination found a follicular hyperplasia in four cases (Fig4-5) and was normal
79 for the remaining case.

80 Four patients experienced an improvement after thymectomy, with a complete remission and
81 a **tolerated medical wean** in case 1 which was AchR-Ab positive. A clinical improvement with
82 medical treatment decrease was obtained in three other patients. **Unfortunately** one patient
83 **could not** be evaluated because of non adherence to medical treatment.

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87 Discussion:

88 Management of JMG was initially adopted from adult patients, whereas recent studies have
89 showed that JMG is different in presentation and in the course of the disease, particularly the
90 group of prepubertal children (3-5).

91 Actually, children with JMG exhibit higher rates of remission than adults. This includes
92 spontaneous remission and remission following a period of drug therapy. Prepubertal children
93 have the highest rates of spontaneous remission (2, 6).

94 Therapeutic strategies for JMG have been established. Medical treatment as first line therapy
95 consists of pyridostigmine associated to steroids and long term immunosuppression in
96 generalised muscular weakness and moderate to severe bulbar symptoms or respiration
97 insufficiency. Plasmapheresis can be indicated in severe forms (1).

98 All of our patients received pyridostigmine, associated to steroids in two cases and steroids
99 with immunosuppression in one case.

100 Because of the presumed role of the thymus in the pathogenesis of myasthenia gravis,
101 thymectomy is a recognised aspect of management. Thymectomy may remove germinal
102 centres and disrupt antibody diversification (2, 7). Childhood thymectomy leads to a
103 premature immunosenescence, mimicking changes expected after physiological thymus
104 involution in the elderly with loss of thymic function (8). This could explain why many
105 researchers have argued against the surgical intervention in prepubertal patients (9).

106 We compared our findings with other series of thymectomised JMG (table2).

107 More patients with thymic hyperplasia achieved remission compared to patients with normal
108 thymus (2). In our series, four patients had thymic hyperplasia, one of them showed a
109 complete remission after surgery.

In JMG thymic hyperplasia is evident in 83% of patients and in 3.8% thymoma could be detected (1, 10).

Although the outcome of thymectomy has not been evaluated with randomised controlled trials in children, some case series have suggested that remission rate following thymectomy in JMG is higher than the remission rate with or without medical treatment (3, 11,12).

Thymectomy is followed by improvement in most cases: remission rate was higher in children after thymectomy than in the group of spontaneous evolution (1, 7, 11, 13-15).

Chao Cheng et al made a comparison of major series of patients with JMG who underwent thymectomy from different areas of the world with his own study (135 patients). He noted a comparable complete remission rate after surgery across the different series, which varied between 37.5 - 60% (12).

Thymectomy is recommended as early as possible in case of generalised weakness (1).

The duration between disease onset and thymectomy is a significant predictor of the efficacy after surgery in generalised JMG. Remission is higher if thymectomy is performed within the first year after onset (1, 6, 12, 14, 16, 17,12).

More recent review of children, including prepubertal patients, also suggested increased remission rates after thymectomy (2, 14, 16, 18).

Chao Cheng et al suggested an algorithm for treatment of juvenile myasthenia gravis (JMG) patients based on age (>12 years old), Osserman classification (>I), duration of the disease (≥ 24 months) and medical treatment response (12).

Current evidence suggests that thymectomy should not be indicated in Musk-positive disease as it is unclear whether it confers any benefit (3, 19-21). In our patients Musk-antibodies were not practiced.

Thymectomy in pure OMG remains controversial. Whereas OMG is not life threatening, patients may be dependent on long term immunosuppressant medications, including corticosteroids with the resultant side effects which can be substantial in children. Thymectomy has been performed in refractory cases (2).

A variety of surgical methods for thymectomy have been described: full or partial sternotomy, thoracoscopic or transcervical approaches (2, 22-23).

Many authors recommend a transsternal approach in children to prevent incomplete removal of all thymic tissue, which may lead to poor outcome (3, 24).

Less invasive techniques such as VAT thymectomy are now resulting in comparable remission rates following thymectomy in adult MG (3, 25).

Della Marina et al recommend thoracoscopic techniques but these are restricted to specialised centers (1).

Kolski HK et al (26) has applied VAT to a group of juvenile patients for thymectomy and compared it to a similar group of six patients (in terms of age and clinical severity) operated via a median transsternal approach. He concluded that VAT thymectomies are comparably effective to transsternal procedures in treating generalised JMG and can be safely performed in children as young as 20 months of age. In addition, VAT surgery is less invasive and significantly shortens the postoperative hospital stay, and has superior cosmetic results.

In our series, thymectomy was performed via antero-lateral thoracotomy in three cases. Thoracotomy was practiced by our surgical team because they mastered it better than sternotomy. VAT was introduced recently and served to treat the two other cases.

A larger series with a randomised controlled study is necessary to elucidate the benefit of thymectomy in JMG and to compare results of different surgical methods.

Conclusion:

The benefit of thymectomy in children with JMG is still controversial. However, many researchers have reported a sustained improvement of symptoms in the majority of patients after thymectomy. Although the number of our patients is too few to indicate any treatment recommendations, our findings seem to encourage thymectomy in case of JMG and particularly when imaging anomalies are found. VAT thymectomy was well tolerated and it is considered by many researchers to be the standard method for thymectomy in children.

Randomised controlled trials are necessary to elucidate the advantages of thymectomy and to establish a clear treatment algorithm for children with JMG.

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Figure legends:

Fig1: Chest x-ray: Mediastinal enlargement.

Fig2: Mediastinal CT-scan: shows the presence of a thymoma.

Fig3: Complete thymectomy.

Fig4: Thymic hyperplasia (H&E, original magnification x200)

Fig5: Thymic hyperplasia (H&E, original magnification x400)

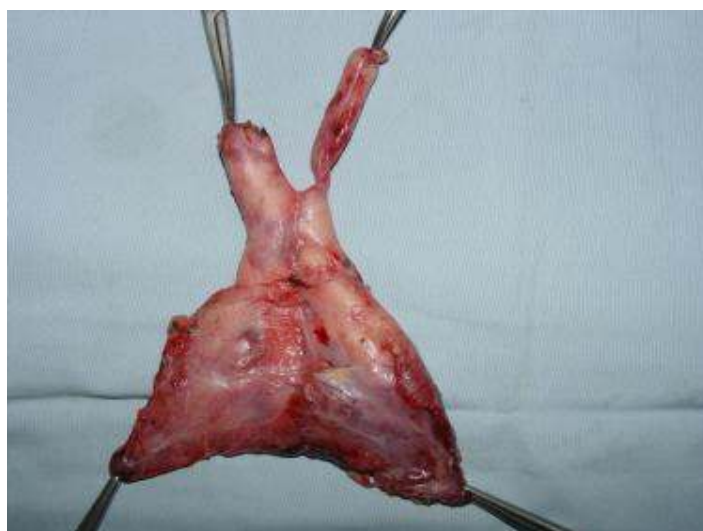


259 Fig1. Chest x-ray



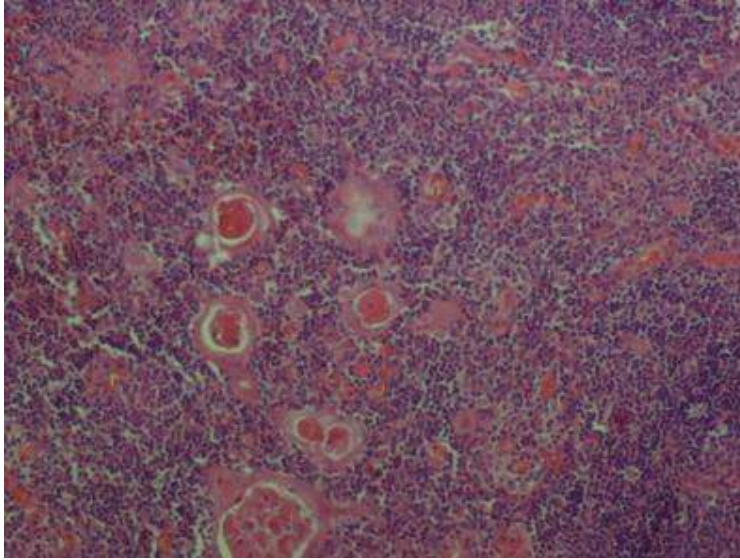
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261 Fig2. Thoracic CT Scan



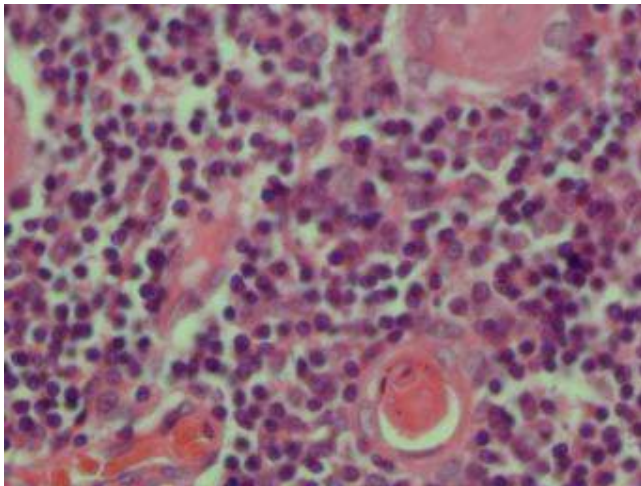
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263 Fig3. Thymectomy



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265 Fig.4 Histology. Thymic hyperplasia (H&E, original magnification x200)



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267 Fig.5 Histology: Thymic hyperplasia (H&E, original magnification x400)

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Table 1: data about the five patients with JMG

Case	Age (year)	Sex	Osseman classification	AchR-Ab	Imaging CT-Scan+/- MRI	Medical treatment Tablets/day	Surgery	Histology	Evolution/ Medical treatment Tablets/day
1	14	M	IIb	Positive	Thymoma	Pyridostigmine 6	Thoracotomy	Hyperplasia	Remission/No
2	2.5	F	IIa	Negative	Hypertrophy	Pyridostigmine 6	Thoracotomy	Hyperplasia	Improvement /yes 3
3	9	F	IIa	Negative	Thymoma	Pyridostigmine 3	Thoracotomy	Hyperplasia	Improvement/yes 2
4	10	F	IIb	Negative	Thymoma	Pyridostigmine 4	VAT	Normal	No improvement+ Non adherence to medical treatment
5	3	F	I	Negative	Hypertrophy	Pyridostigmine 3	VAT	Hyperplasia	Improvement/yes 2