The feasibility of quantitative MRI of extra-ocular muscles in myasthenia gravis and Graves' orbitopathy

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Although quantitative MRI can be instrumental in the diagnosis and assessment of disease progression in orbital diseases involving the extra-ocular muscles (EOM), acquisition can be challenging as EOM are small and prone to eye-motion artefacts. We explored the feasibility of assessing fat fractions (FF), muscle volumes and water T2 (T2water) of EOM in healthy controls (HC), myasthenia gravis (MG) and Graves' orbitopathy (GO) patients. FF, EOM volumes and T2water values were determined in 12 HC (aged 22-65 years), 11 MG (aged 28-71 years) and six GO (aged 28-64 years) patients at 7 T using Dixon and multi-echo spin-echo sequences. The EOM were semi-automatically 3D-segmented by two independent observers. MANOVA and t-tests were used to assess differences in FF, T2water and volume of EOM between groups (P < .05). Bland–Altman limits of agreement (LoA) were used to assess the reproducibility of segmentations and Dixon scans. The scans were well tolerated by all subjects. The bias in FF between the repeated Dixon scans was −0.7% (LoA: ±2.1%) for the different observers; the bias in FF was −0.3% (LoA: ±2.8%) and 0.03 cm3 (LoA: ± 0.36 cm3) for volume. Mean FF of EOM in MG (14.1% ± 1.6%) was higher than in HC (10.4% ± 2.5%). Mean muscle volume was higher in both GO (1.2 ± 0.4 cm3) and MG (0.8 ± 0.2 cm3) compared with HC (0.6 ± 0.2 cm3). The average T2water for all EOM was 24.6 ± 4.0 ms for HC, 24.0 ± 4.7 ms for MG patients and 27.4 ± 4.2 ms for the GO patient. Quantitative MRI at 7 T is feasible for measuring FF and muscle volumes of EOM in HC, MG and GO patients. The measured T2water was on average comparable with skeletal muscle, although with higher variation between subjects. The increased FF in the EOM in MG patients suggests that EOM involvement in MG is accompanied by fat replacement. The unexpected EOM volume increase in MG may provide novel insights into underlying pathophysiological processes.

Abbreviations used: AChR, acetylcholine receptor; EOM, extra-ocular muscles; EPG, extended phase graphs; FF, fat fraction; GO, Graves’ orbitopathy; HMS, hereditary motor and sensory neuropathy; IR, inferior rectus muscle; LP, levator palpebrae muscle; LR, lateral rectus muscle; MG, myasthenia gravis; MR, medial rectus muscle; MSE, multi-echo spin-echo; MuSK, muscle-specific tyrosine kinase; NO, optic nerve; SR, superior rectus muscle; SR+LP, superior rectus and levator palpebrae muscle complex; TM, temporalis muscle; TSH-R, thyroid-stimulating hormone receptor.

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1 | INTRODUCTION

In diseases that involve the extra-ocular muscles (EOM), local muscle weakness or muscle swelling results in complaints such as double vision and ptosis. The auto-immune disease myasthenia gravis (MG) is characterized by fatigable muscles. Auto-antibodies, targeting the acetylcholine receptor (AChR) or muscle-specific tyrosine kinase (MuSK), disturb transmission at the neuromuscular junction. In 85% of patients, MG starts with diplopia and ptosis, which are both caused by EOM weakness. Currently, MG is diagnosed by either the presence of auto-antibodies in serum, or abnormal findings during neurophysiological testing. However, in patients with pure ocular symptoms, the sensitivity of these tests is low (0.44 for testing antibodies, 0.34 for repetitive nerve stimulation and variable between 0.66 and 0.98 for single-fiber electromyography). Consequently, diagnosis is challenging and more sensitive diagnostic tests are required. Moreover, there is a need for prognostic markers to predict disease progression in ocular MG and refractory MG.

Graves’ orbitopathy (GO) is an orbital disease caused by auto-antibodies acting against thyroid proteins, including the thyroid-stimulating hormone receptor (TSH-R). Due to inflammation and swelling of the EOM, symptoms of GO include upper eyelid retraction, proptosis and diplopia. At a later stage of the disease, fibrosis appears in the EOM as a result of the longstanding inflammation. There is an unmet clinical need to categorize patients, since immunosuppressant medication and radiotherapy are only effective in the active stage of GO.

Quantitative MRI can be used to assess disease progression and to study the pathophysiology of skeletal muscle in neuromuscular disease by assessing fat replacement, muscle size and T2 relaxation time changes. Assessment of such structural changes in individual EOM by quantitative MRI may therefore also contribute to understanding the pathophysiology and pattern of muscle involvement of the EOM in orbital disease. As such, it could contribute to identifying diagnostically challenging subgroups in GO and aid in the follow-up of therapeutic response in MG.

In the current study we explored the feasibility and value of quantitative MRI of the EOM in MG and GO patients and healthy controls. These specific patient groups were chosen because distinct structural changes are expected in both diseases and additional diagnostic tests could directly improve clinical care. In MG there is histological evidence for fat replacement, fibrosis and atrophy of the EOM, while in GO the EOM are known to be swollen and inflamed at an early stage, and fat-replaced and fibrotic at a later stage of the disease. However, the EOM are challenging to assess with quantitative MRI. Since they are small, the high resolution needed for MRI of these muscles results in a poor signal-to-noise ratio (SNR). In addition, MRI of the EOM is prone to eye-motion artefacts, which reduce the image quality. Moreover, the surrounding bone from the orbit and air from the nasal sinuses results in an inhomogeneous B0 field. However, recent studies have shown that the increased SNR at 7 T, combined with a cued blinking paradigm to reduce eye-motion artefacts and localized shimming to minimize B0 artefacts, enables high resolution imaging of the eyes. Based on these protocols we therefore developed a dedicated protocol for anatomical and functional imaging of the EOM.

2 | METHODS

2.1 | Participants

All patients were recruited from the outpatient clinics of the neurology and the ophthalmology departments of the Leiden University Medical Center. The diagnosis of MG was confirmed by a neurologist, based on a combination of clinically confirmed fluctuating muscle weakness and the presence of AChR antibodies or abnormal decrement by repetitive nerve stimulation. The diagnosis of GO was confirmed by an ophthalmologist, based on the presence of the characteristic orbital abnormalities (eyelid retraction, redness/swelling of the eyelids/conjunctiva and enlarged EOM or orbital fat) and the presence of anti-TSH-R antibodies. For MG and GO patients, both acute (diagnosis less than 3 months ago) and chronic (diagnosis more than 1 year ago) stages of the disease were included, to assess a wide variety of structural changes to study the sensitivity of the dedicated MRI protocol. Healthy controls were recruited from the radiology database for healthy controls.

The local medical ethics committee approved the study and all study participants gave written informed consent prior to MR scanning.

2.2 | MR examination

We performed chemical shift-based water fat separation using the Dixon method to quantify fat replacement and muscle volume and multi-echo spin-echo (MSE) to assess the T2 relaxation time of water (T2water) as an indicator of disease activity. All subjects were scanned in supine
position on a 7 T Philips Achieva MRI (Philips Healthcare, Best, the Netherlands) using a cued blinking paradigm\(^3\) with the upper 16 elements of a 32-channel head-coil (Nova Medical). Patients were asked to blink upon visual cues provided on a screen; the MRI acquisition was halted during blinking. A three-point multi-acquisition 3D Dixon scan was acquired (resolution: 0.8 x 0.8 x 0.8 mm\(^3\), first time to echo [TE] = ΔTE/repetition time [TR]/flip angle [FA]/scan time: 2.4 ms/0.33 ms/10 ms/3/3 minutes 50 seconds). The 3D volumetric Dixon scan was planned with the frequency encoding in the left-right direction and was angulated in the coronal plane if necessary, to include both orbits in the field of view. In a subset of seven MG patients and five healthy controls, a second Dixon scan was acquired (resolution: 0.7 x 1.0 x 0.7 mm\(^3\), first TE/ΔTE/TR/FA/scan time: 2.4 ms/0.33 ms/10 ms/3/3 minutes 50 seconds) to study the value of a higher resolution in the coronal plane, while maintaining the same voxel volume. These Dixon scans with a higher resolution in the coronal plane were used to study the reproducibility of the fat fraction (FF) measurement by comparing them with the isotopic Dixon scan. In another subset of participants, containing five MG patients, one GO patient and seven healthy controls, a MSE scan was acquired per orbit (resolution: 1.2 x 1.2 x 3.0 mm\(^3\), first TE/ΔTE/TR: 12 ms/12 ms/4000 ms, 24 echoes, three slices per eye, scan time: 2 minutes 44 seconds). This coronal scan was planned with one orbit in the field of view perpendicular to the medial rectus muscle (MR) and lateral rectus muscle (LR). The scan times were kept under 4 minutes, as previous experience shows this minimizes movement artefacts. Higher order shimming was performed for all scans, to minimize the susceptibility artefacts caused by the bony structures of the orbit and the nasal cavities.

### 2.3 Data analysis and segmentation

Dixon scans were reconstructed with seven-peak reconstruction using the manufacturer’s software to yield water and fat images. No T2\(^*\) correction was performed, given the short echo spacing of 0.33 ms. To minimize eye movement artefacts, the scan time was minimized by using a relatively low TR of 10 ms. Correction for the resulting T1-weighting in the water and fat images was incorporated in the FF calculation, assuming T1 values of 1400 ms (correction factor: 1.22) and 365 ms (correction factor: 1.08) for water and fat, respectively\(^21\):

\[
\text{FF} = \frac{1.08 \times \text{Fat}}{1.22 \times \text{Water} + 1.08 \times \text{Fat}}
\]

Scans were excluded if there were obvious motion artifacts. For the Dixon scans, the four rectus EOM (LR, MR, inferior rectus muscle [IR] and the superior rectus and levator palpebrae muscle complex [SR+LP]) were semi-automatically 3D-segmented on the water image, using a seed-growing algorithm with a manually set threshold for each subject in ITK-SNAP\(^22\) (Figure 1) by one observer (KRK). After automatic segmentation, the superior rectus and levator palpebrae muscle complex [SR+LP] were semi-automatically 3D-segmented on the water image, using a seed-growing model. This is especially important at high field strengths, as the shift scales linearly with the magnetic field strength and is 1.2 mm for the direction between the excitation and refocusing pulses, due to different slice selection gradients for both pulses, was also incorporated in the model. This is especially important at high field strengths, as the shift scales linearly with the magnetic field strength and is 1.2 mm for the performed MSE scan. A dictionary-matching algorithm was used to obtain the T2 of the water component, the B1 map and FF for each voxel.\(^{23,25}\) This dictionary was created using T2\(_{\text{water}}\) values from 10 to 60 ms, T2\(_{\text{fat}}\) values from 120 to 200 ms, FF values from 0% to 100% and B1 values from 50% to 140%. To obtain a T2\(_{\text{fat}}\) value for the match, the T2\(_{\text{fat}}\) is conventionally calibrated on the subcutaneous fat, assuming it consists only of fat tissue and has a similar T2 as the fat replacement in skeletal muscles.\(^{26}\) However, in this work the T2\(_{\text{fat}}\) was calibrated on the orbital fat, since subcutaneous fat was not present in the field of view. For the MSE scans, the EOM were segmented by drawing ROIs per slice using the FF map as an anatomical reference by one observer (KRK).

### 2.4 Statistics

The Bland–Altman bias and limits of agreement between the FFs of the two Dixon scans were calculated for all participants together, and separately for the healthy controls and MG patients. The FFs and muscle volumes from the segmentation performed by the two independent
observers were compared using Bland–Altman analysis. Mean FF and muscle volumes were compared using MANOVA with post hoc tests with Bonferroni multiple-testing correction on the patient average and on the individual muscles after averaging the EOM of the left and right eye per patient. Mean $T_{2\text{water}}$ values were compared using a two-tailed t-test between healthy controls and MG patients, but not for the GO patient, since only one observation was present. The correlation between age and FF was tested using Pearson correlations for all three groups separately. The correlation between FF and volume was tested for healthy controls for all EOM. Significance level for all tests was set at .05.

3 | RESULTS

3.1 | Subject characteristics

Twelve healthy subjects (aged 41.0 ± 15.9 years, 42% male), eleven MG patients (aged 52.5 ± 15.9 years, 64% male) and six GO patients with active disease (aged 47.7 ± 14.0 years, 50% male, clinical activity score $\geq 3$) were included in this pilot study (Table 1).

3.2 | Data quality

All scans, except for one Dixon scan with higher resolution in the coronal plane from an MG patient, were free from visible motion artefacts. This scan was excluded from the analysis. Reconstructed water, FF and $T_{2\text{water}}$ maps and the first and last echo of the MSE scan are shown in Figure 2.

3.3 | Agreement on semi-automatic segmentation

The EOM were semi-automatically segmented by two independent observers separately. Figure 3 shows Bland–Altman plots depicting the bias and limits of agreement between the two independent observers for FF (A) and volume (B). The mean bias and limits of agreement in FFs and volume were $-0.3\%$ ([95% CI]: $-3.1\%; 2.6\%$) and 0.03 cm$^3$ ([95% CI]: $-0.33$ cm$^3; 0.39$ cm$^3$), respectively.

3.4 | Reproducibility of FF measurement by Dixon scans

In one patient the second Dixon scan was excluded from the analysis due to severe motion artefacts. On visual inspection, the isotropic Dixon scan was of a higher quality compared with the Dixon scan with a higher resolution in the coronal plane. The gain of resolution in the coronal plane...
plane was accompanied by a loss of resolution in the transversal plane to maintain sufficient SNR and therefore unwanted partial volume effects were present. The Bland–Altman bias and limits of agreement between the two Dixon scans were $-0.7\%$ (95% CI: $-3.4\%$; $1.8\%$) for all participants, with the Dixon scan with a higher resolution in the coronal plane giving, on average, lower FFs. In the MG patient group the bias and limits of agreement were $-1.0\%$ (95% CI: $-3.7\%$; $1.7\%$), and in healthy controls the level of agreement was $-0.4\%$ (95% CI: $-2.6\%$; $1.7\%$) (Figure 3C).

### 3.5 | Muscle volume

The mean muscle volumes of the four recti muscles were higher in GO patients ($1.2 \pm 0.4 \text{ cm}^3$) than in healthy controls ($0.6 \pm 0.2 \text{ cm}^3$; $P < .001$) (Figure 4). The mean EOM volume was also higher in MG patients ($0.8 \pm 0.2 \text{ cm}^3$) than in healthy controls ($P = .007$). Assessing all four muscles

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| TABLE 1 | Baseline characteristics of the three groups |
|---------------|--------------------------------------------------|
|               | Healthy control (n = 12) | Myasthenia gravis (n = 11) | Graves' orbitopathy (n = 6) |
| Age, years    | $41.0 \pm 15.9$          | $52.5 \pm 15.9$           | $46.7 \pm 14.0$          |
| Gender        |                         |                         |                           |
| Male          | 5 (42%)                 | 7 (64%)                 | 3 (50%)                  |
| Female        | 7 (58%)                 | 4 (36%)                 | 3 (50%)                  |
| Disease onset |                         |                         |                           |
| <3 months     | -                       | 5 (45%)                 | 3 (50%)                  |
| >1 year       | -                       | 6 (55%)                 | 3 (50%)                  |
| Medication    |                         |                         |                           |
| Pyridostigmine| -                       | 10 (91%)                | -                        |
| Prednisone    | -                       | 6 (55%)                 | 2 (33%)                  |
| Other immunosuppressants | -          | 1 (9%)                 | -                        |
| MG phenotype  |                         |                         |                           |
| Ocular        | -                       | 7 (64%)                 | -                        |
| Generalized   | -                       | 4 (36%)                 | -                        |
| Anti-TSH-R antibodies | -          | -                      | 6 (100%)                 |
| Anti-AChR antibodies | -          | 10 (91%)               | -                        |

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**FIGURE 2** Examples of water images and fat fraction maps of the isotopic Dixon scan and the Dixon scan with a higher resolution in the coronal plane on the left. On the right, the first echo, the last echo and the $T_{2\text{water}}$ map are shown for the multi-echo spin-echo scan. The colored arrows point out the lateral rectus muscle (LR; red), the medial rectus muscle (MR; green), the inferior rectus muscle (IR; blue) and the superior rectus and levator palpebrae muscle complex (SR+LP; yellow)
separately, the volumes of all four rectus EOM differed significantly between GO patients and healthy controls (LR: \( P = .033 \); MR: \( P = .001 \); IR: \( P < .001 \); and SR+LP: \( P = .001 \)). No significant differences were observed comparing individual EOM between healthy controls and MG patients. A difference between GO and MG patients was present in the MR (\( P = .035 \)) and the IR (\( P = .001 \)) (Figure 5).

### 3.6 | FFs

The mean FFs of the EOM were higher in MG patients (14.1% ± 1.6%) compared with those of healthy controls (10.4% ± 2.5%; \( P = .008 \)) (Figure 6) and similar between MG and GO patients (13.9% ± 4.4%). The variability in FFs appeared higher in GO patients than in MG patients and healthy controls. When analyzed individually, all four rectus EOM differed between MG patients and healthy controls (LR: \( P = .011 \); MR: \( P = .048 \); IR: \( P < .003 \); and SR+LP: \( P = .003 \)). A difference between healthy controls and GO patients was observed in the LR (\( P = .029 \)) and the SR+LP (\( P = .003 \)) (Figure 7). For each group, Pearson correlations did not indicate a relationship between FF and age (\( r = 0.006 \) and \( P = .86 \) for healthy controls; \( r = 0.21 \) and \( P = .69 \) for GO patients; and \( r = 0.31 \) and \( P = .38 \) for MG patients). No dependence of FF on volume was observed in healthy controls (slope: −2.1%/cm³ [95% CI: −4.3 to 0.1]; \( r = 0.19 \) and \( P = .06 \)).

### 3.7 | Water T2

The average \( T_2 \) for all EOM was 24.6 ± 4.0 ms for healthy controls, 24.0 ± 4.7 ms for MG patients and 27.4 ± 4.2 ms for the GO patient. No significant differences between healthy controls and MG patients were found. \( T_2 \) values for individual EOM are depicted in Figure 8.
DISCUSSION

The current study aimed to study the feasibility of quantitative MRI of EOM in diseases that affect the eye muscles. We showed that the EOM could be reproducibly segmented semi-automatically with a small inter-observer variation and that FFs could be reproducibly measured in healthy...
controls and MG and GO patients between different scans. Thus we were able to detect small differences in muscle volume and FF between healthy controls and two patient cohorts, as well as $T_2^{\text{water}}$ values similar to spectroscopy-measured values in skeletal muscles, although small differences cannot be excluded as the study was not powered to detect them.

All scans except one were free from visible motion artefacts, indicating that all subjects could adhere to the protocol. The mean bias in FF between the segmentations of two independent observers was low (0.3%) and the limits of agreement were within ±3% in FF, which is in the same range as previously reported values in segmentation of the vastus lateralis in patients with Duchenne muscular dystrophy (bias: 0.1%, limits of agreement: ±1.1%), and in patients with myotonic dystrophy type 1 in lower leg muscles (limits of agreement: ±2.6%). The scan-to-scan reproducibility between the two Dixon scans with different resolutions showed a low average bias in FF (−0.7%) and narrow limits of agreement (−3.4% to 1.8%), which is lower than the difference found between the patients and the healthy controls (ie, ~4%), indicating the feasibility of assessing FFs using this method to detect affected muscles. For comparison, Dixon measurements in skeletal muscles have been reported to have a reproducibility (limits of agreements) of ±1% for water-fat phantoms, ±1.1% for thigh muscles in healthy controls, and ±2.5% for the leg muscles in patients with Duchenne muscular dystrophy. These techniques could be used in future longitudinal studies to study the disease progression in conditions affecting the EOM and to relate time-varying clinical observations such as painful eyes to the current condition of the EOM in terms of fat replacement and inflammation.

We found FFs of ~10% in the EOM, which is higher than the 2%-5% commonly observed in skeletal muscles of healthy volunteers. Although the high FF could be an intrinsic property of EOM, it could also be the result of noise bias. Earlier studies in skeletal muscles have shown that low SNR could result in an overestimation of the FF in the low FF ranges. Although the strong contrast between the orbital fat and the muscles and globe suggests more than sufficient SNR for anatomical evaluation of the images, some noise can be observed in the temporalis muscle (TM), a large skeletal muscle close to the orbit. A Dixon reconstruction of 4-fold downsampled source images of the same subject, which increased the SNR 2-fold, resulted in a reduction in FF from 9.3% to 5.1%, showing that the protocol used is susceptible to noise bias in the low FF ranges. This could explain the relatively high FF observed in the EOM. Therefore, we recommend that future studies use a higher SNR for the Dixon acquisition, for example, by using a higher FA or a two FA approach. Additionally, the overestimation of the FF could have been caused by partial volume effects of the EOM with the surrounding orbital fat, due to the small size of these muscles (~1 cm$^3$).

Muscle volumes were measured using 3D segmentation on the water image from the Dixon scans. The limits of agreement for the muscle volumes segmented by the two independent observers ranged between −0.33 and 0.39 cm$^3$. These limits do not reach the 0.6 cm$^3$ difference observed between GO patients and healthy controls, but are higher than the difference of 0.2 cm$^3$ observed between the MG patients and healthy controls. On visual inspection (Figure 3), the highest differences between observers were found in the SR+LP, which is the hardest to delineate because the complex contains two separate muscles. An overall dependence of volume difference between observers on mean volume was observed (Figure 3A), but there was virtually no mean bias (0.03 cm$^3$). Moreover, as no dependence of FF on the volume was present in healthy controls, the segmentation was not likely to have influenced the observed FF differences between groups.

Using MSE and EPG analysis, we found $T_2$ values of the myocytic compartment ($T_2^{\text{water}}$) for EOM to be comparable with the $T_2^{\text{water}}$ in healthy skeletal muscle, as measured with MR spectroscopy. The standard deviation of $T_2^{\text{water}}$ between the EOM (4.0, 4.7 and 4.2 ms for healthy controls, MG patients and GO patients, respectively) was higher than previously measured using an EPG model in leg muscles.
between subjects (1.4-2.1 ms). This variation between T2\textsubscript{water} of muscles was expected as different patients have different patterns of muscle involvement. For example, in GO the IR is usually affected earlier in the disease, followed later by the other EOM. In the healthy controls, however, a smaller variation, comparable with other skeletal muscles, was expected. Two assumptions in the analysis may have contributed to this higher variation. Firstly, the calibration of the T2\textsubscript{fat} was performed on the orbital fat, under the assumption that its composition is similar to intra-muscular fat. Orbital fat is, however, known to be involved in inflammatory conditions, such as GO. This variation in orbital fat T2, which is not necessarily also present in the T2 of intra-muscular fat, might result in an erroneous variation in intra-muscular T2\textsubscript{fat}. Secondly, our model assumes that the ROI to calibrate the T2\textsubscript{fat} only contains orbital fat. However, in reality there is vascularization in the orbital fat that will result in a small, patient-specific inaccuracy in the measured orbital T2\textsubscript{fat}, which will subsequently influence the obtained T2\textsubscript{water}. A high variation between individual EOM in patients is to be expected, as MG and GO are known to selectively affect specific EOM. Due to the high variation, the power was limited to detect mean differences in T2\textsubscript{water} between groups. For future studies, we recommend establishing baseline values per EOM and using Z-scores to detect EOM with an abnormal T2\textsubscript{water}.

Although the patient cohorts were small in size, clear differences were found between them. The measured FFs of the EOM were higher in MG and GO patients compared with healthy controls and muscle volumes were higher in GO patients. Edematous EOM are a common finding in GO patients and fat replacement of EOM has been described in late-stage GO patients. The variability in muscle volume was high in GO patients, possibly due to the inclusion of patients with varying disease severity and the inflammation of only a portion of the EOM in individual patients.

The increase in FF in the EOM in MG has been described in histological studies, but the increase in muscle volume in the EOM of MG patients was unexpected. In MG, AChR or MuSK, antibodies can result in structural damage of the muscle membrane at the neuromuscular synapse, which is usually located at the middle of the muscle fiber, but this is not a dystrophic process. In dystrophic muscle diseases fat replacement of muscle tissue is commonly observed, and can almost completely replace normal muscle tissue. In neuropathic diseases, like hereditary motor and sensory neuropathy (HMSN) or hereditary neuropathy with liability to pressure palsies, FFs of up to 15% have been observed in calf muscle, which represents a relatively small increase (of up to 10%) in FF, and is comparable with the increase in FF (of up to 8%) that we observed in the EOM in MG. In a recent case report, histological findings in a seronegative MG patient also included adipocyte replacement of muscle fiber, and fat replacement has been seen in the bulbar muscles of MuSK-positive MG patients. Our results agree with these observations and indicate that involvement of EOM in MG patients is accompanied by fat replacement. Interestingly, our results to date suggest that the FFs of MG patients with a disease onset of less than 3 months before the scan did not appear to differ from those of chronic MG patients.

Severe MG can induce chronic denervation and therefore might behave as a neuropathic condition. Muscular atrophy in skeletal muscle is observed in 10% of AChR-positive MG patients with longstanding disease. In MuSK-positive MG patients, early atrophy is seen in the bulbar muscles. In other neurogenic muscle diseases, like HMSN, a decrease in muscle volume has also been observed. However, in this work we observed an increase in EOM volume in MG patients, which is not consistent with the expected atrophy. This unexpected increase in muscle volume raises questions about its pathophysiology. Firstly, it is possible that inflammatory processes in the EOM result in swollen muscles. In MG, complement activation is observed, resulting in activation of the membrane attack complex and destruction of the neuromuscular endplate. To what extent this inflammatory process affects the EOM is unknown. Secondly, (pseudo)hypertrophy may provide an explanation of the increase in EOM volume that has been observed in some muscular dystrophies.

A limitation of this work was that the groups in this pilot study were not fully age-matched, as the maximum age of healthy volunteers was limited to 65 years, based on the inclusion criteria of our institutional review board approval. This upper age limit did not apply to both patient groups. However, an earlier study showed that the EOM do not change in muscle thickness, size and fatty tissue volume between age groups. In addition, when analyzing all three groups in this study separately, we did not see any effects of age on EOM FFs. Therefore, age does not appear to be a major confounder in our study. Secondly, due to the exploratory nature of this pilot study with ongoing sequence development, not all sequences were applied to all patients. Consequently, the second Dixon scan for reproducibility and the MSE sequence were only performed in a subgroup of patients, and T2 relaxometry was only applied to one GO patient. Finally, this study was performed on a 7 T MR scanner to increase the SNR at the high resolution required for imaging the EOM. However, the disadvantages of scanning at 7 T are the increase in susceptibility differences, B0 inhomogeneities and chemical shift artefacts. These challenges were successfully solved using higher order shimming and the in-plane chemical shift displacement was set at 1.0 acquisition pixels and corrected for during postprocessing. Future studies can determine if the same results can be achieved at clinical field strengths such as 3 T.

In conclusion, despite the challenges of size and movement, quantitative MRI is feasible for measuring FFs and muscle volumes of individual EOM in healthy controls, MG and GO patients at 7 T. We recorded water T2 relaxation times that were comparable with values measured in skeletal muscle, but found high variability between individuals and between different EOM. Detection of group differences for this parameter will therefore only be possible when large differences between groups are present, only affected EOM are included in the analyses, or when using large sample sizes. We found an increase in EOM volume and FF in MG and GO patients. The unexpected increase in EOM volume in MG may provide novel insights into underlying pathophysiological processes. The diagnostic value of these findings and their value when following up disease progression should be the subject of future prospective studies.
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DATA AVAILABILITY STATEMENT

The used Matlab scripts and one sample dataset presented in this article will be made available at the request of a qualified investigator. Requests should be made to K. R. Keene (k.r.keene@lumc.nl).

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