Evaluation of the efficacy and safety of MRI-guided focused ultrasound (MRgFUS) for focal hand dystonia: study protocol for an open-label non-randomised clinical trial

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ABSTRACT

Introduction MRI-guided focused ultrasound (MRgFUS) thalamotomy provides an exciting development in the field of minimally invasive stereotactic neurosurgery. Current treatment options for focal hand dystonia are limited, with potentially more effective invasive stereotactic interventions, such as deep brain stimulation or lesioning therapies, rarely used. The advent of minimally invasive brain lesioning provides a potentially safe and effective treatment approach with a recent pilot study establishing MRgFUS V0-complex thalamotomy as an effective treatment option for focal hand dystonia. In this study, we undertake an open-label clinical trial to further establish MRgFUS V0-complex thalamotomy as an effective treatment for focal hand dystonia with greater attention paid to potential motor costs associated with this treatment. To elucidate pathophysiology of dystonia and treatment mechanisms, neurophysiological and MRI analysis will be performed longitudinally to explore the hypothesis that neuroplastic and structural changes that may underlie this treatment benefit.

Methods and analysis A total of 10 participants will be recruited into this open-label clinical trial. All participants will undergo clinical, kinematic, neurophysiological and radiological testing at baseline, followed by repeated measures at predesignated time points post MRgFUS V0-complex thalamotomy. Further, to identify any underlying structural or neurophysiological abnormalities present in individuals with focal hand dystonia, 10 age and gender matched control participants will be recruited to undergo comparative investigation. These results will be compared with the intervention participants both at baseline and at 12 months to assess for normalisation of these abnormalities, if present.

Ethics and dissemination This trial was reviewed and approved by the St Vincent’s Health Network Sydney Human Research Ethics Committee (2022/ETH00778). Study results will be published in peer-reviewed journals and presented at both national and international conferences.

Trial registration number CTRN12622000775718.

INTRODUCTION

Dystonias are a complex group of movement disorders that are characterised by sustained or intermittent muscle contractions resulting in abnormal movements, postures or both. Focal dystonia refers to those dystonias where the abnormal movement is isolated to a single body part. Involvement of the upper limb is termed focal hand dystonia and can be classified as either a task-specific action dystonia, whereby the dystonia can be predictably triggered by performance of a specific motor task, or paroxysmal, with episodes of dystonia that occur without specific task performance or persistent. For the purposes of this manuscript, focal hand dystonia refers to that...
Current treatment options for focal dystonia are limited to short term symptom relief with low efficacy oral medications or local treatment with botulinum toxin. Physical therapies and rehabilitation have been reported to provide some improvement, but this evidence is not definitive. Unlike generalised or segmental dystonia syndromes, where the GPi (globus pallidus internus) is preferred, small non-randomised trials suggest that focal hand dystonia responds well to targeted intervention of the thalamus; specifically, the Vo complex. Thalamic deep brain stimulation (DBS) is an effective treatment option for individuals with writer’s cramp; however, there are inherent risks associated with DBS, including perioperative anaesthetic complications, intraparenchymal haemorrhage, electrode/lead breakage or infection. The use of stereotactic thalamotomy, which involves an open surgical or radiofrequency incision of the thalamus, has demonstrated significant improvement in focal hand dystonia but is similarly an invasive treatment modality. Gamma-knife ablation of the Vo-complex has reported positive outcomes in musician’s dystonia, however, this carries the risk of radiation exposure and neoplasia. MRI-guided focused ultrasound (MRgFUS) is a minimally invasive functional neurosurgical technique that aims to avoid the risk associated with more invasive stereotactic surgical modalities. This technique involves the administration of targeted beams of ultrasound energy through the skull, without requiring a surgical incision, to ablate specific neural pathways without damaging the surrounding brain tissue. MRgFUS thalamotomy is a safe and effective treatment option for multiple tremor syndromes and has both Therapeutic Goods Administration (TGA) and US Food and Drug Administration approval for the treatment of essential tremor. The use of MRgFUS Vo thalamotomy for focal hand dystonia, without tremor, was proposed as an alternative to more invasive stereotactic surgery in 2018, with the first case report appearing soon after. Since this time, a recent pilot study has provided evidence that MRgFUS thalamotomy provides relief of the symptoms of focal hand dystonia with clinical benefit extending out to 12 months post-treatment. While the thalamus has an important role in motor planning and execution, it also has a complex role in the integration and processing of non-motor information, including sensory perception, arousal modulation, limbic regulation of mood and motivation, and associative inputs connoting cognition. Thalamic anatomy can be parcellated based on neuropathological and functional neuroimaging to allow the targeted intervention of motor pathways with preservation of non-motor thalamic function, however, there is little data reporting the loss of fine motor control, learning or task imprecision following MRgFUS. It may be that,

dystonia which is task-specific and isolated to the upper limb. The most common subtypes of focal task-specific hand dystonia are ‘musician’s dystonia’ and ‘writer’s cramp’. Musician’s dystonia presents as muscular incoordination, or loss of voluntary control, in extensively trained musicians. The specific pattern of abnormal ordination, or loss of voluntary control, in extensively trained musicians is significantly higher in those aged 50 and over. Similarly, writer’s cramp is characterised by involuntary movements and posturing that typically affects the fingers, hand and forearm and occurs during writing.

Estimates of the prevalence of dystonia vary depending on the subtype and the population studied. General estimates suggest that approximately 16–30 per 100,000 individuals are affected, however the incidence of idiopathic dystonia is significantly higher in those aged 50 years or older (732 per 100,000). While dystonia may be more common among an ageing population, there is marked variability in prevalence due to a lack of clear clinical criteria and a wide range of symptom severity. Focal hand dystonia, specifically, affects approximately 7–70 individuals per million in the general population, with a higher prevalence reported in at-risk populations, such as professional musicians where dystonia has been noted in up to 14% of musician-related health clinic presentations.

Individuals with focal dystonia are often disabled and unable to perform the task related to their dystonia with subsequent implications on their profession, livelihood and quality of life. The aetiology of focal hand dystonia is unknown, however, both genetic and environmental factors have been implicated. Individuals with focal dystonia have demonstrated abnormalities in sensory processing, sensorimotor organisation, motor excitability and increased cortical plasticity. It is proposed that, while it may speed motor learning, this increased cortical plasticity may predispose to the formation of unwanted motor associations, hence task specific involuntary movements. Recent investigation suggests that focal dystonia is likely a manifestation of abnormalities of the entire sensorimotor network that is amenable to targeted neurosurgical approaches.

The information obtained from this trial may further establish MRgFUS thalamotomy as an effective treatment option for individuals with focal hand dystonia. However, a significant portion of informed decision-making relates to the potential motor cost that may occur as a result of the thalamotomy. It may be that, while MRgFUS thalamotomy reduces the symptoms of focal hand dystonia, there is slowing and imprecision of movement postprocedure, and this information is crucial in informing both clinicians and patients regarding the best potential treatment option.
while MRgFUS thalamotomy reduces the symptoms of focal dystonia, there is slowing of movement with imprecision, or deautomation, of sequential learnt movements post-thalamotomy. Given the precision, accuracy and tempo required for high level musical performance, either inadequate treatment efficacy or potential motor costs may be significant for professional musicians. The balance of symptomatic benefit and motor cost is crucial in informing joint clinician-patient treatment decisions. Further, the effect of MRgFUS thalamotomy on neuroplasticity and motor learning is yet to be understood.

Study rationale
The advent of MRgFUS thalamotomy in the treatment of focal hand dystonia provides an exciting development in the field of minimally invasive stereotactic neurosurgery. The ventro-oral complex, or Vo-complex, of the motor thalamus acts as the main input structure for pallidal fibres and is an important node in the pallidothalamic network. Pallidal afferents to the Vo-complex assist in the programmed control of movement patterns and are modified with repeated trials of a motor task, indicating the important role of these pathways in motor learning.25 Stereotactic thalamotomy of the Vo-complex is an established method of treatment in focal hand dystonia that provides long-term relief of symptoms.17 26–28 A recent retrospective analysis of 171 patients who underwent surgical Vo-complex thalamotomy for focal hand dystonia demonstrated sustained symptom relief at 12-month follow-up, with recurrence in only 10% of patients.29 MRgFUS thalamotomy provides a minimally invasive incisionless alternative to open surgical thalamotomy and recent case reports have demonstrated safety and feasibility of this approach in focal hand dystonia.22 30 31

Horisawa et al25 published the first open-label trial of MRgFUS thalamotomy in writer’s cramp and musician’s dystonia with sustained improvement in symptoms reported at 12 months. Symptoms of dystonia recurred in 3 patients (30%). This trial established the feasibility of minimally invasive approaches for thalamic lesioning in focal hand dystonia. However, given that task specific focal hand dystonia occurs in the context of performing a highly practised, repeated skill, a pure reduction in the severity of the dystonia, following thalamotomy, may not necessarily equate to a desirable treatment outcome. Further, the procedure may be associated with quantifiable slowing of movement, with imprecision or even deautomation of sequenced learnt movements, post-thalamotomy. This information is crucial in informing treatment decisions prior to consideration of thalamotomy. It is our hypothesis that, while MRgFUS Vo-complex thalamotomy may improve the symptoms of dystonia, as measured by validated questionnaires and task performance, it may result in quantifiable losses in hand strength, velocity and function. To this end, we plan to use validated task specific dystonia questionnaires in addition to objective measures of finger and grip strength, as well as functional upper limb task measurement, both pre-treatment and post-treatment with between group comparisons.

A secondary aim of this study is to assess the effect of MRgFUS Vo-complex thalamotomy on the abnormal synaptic plasticity, cortical inhibition and basal ganglia/cerebellar network changes that are thought to underlie the pathophysiology of focal hand dystonia. Given that Vo-complex thalamotomy has been shown to reduce the symptoms of focal hand dystonia, we hypothesise that disruption of the pallidothalamic tract, via thalamotomy, will lead to changes in neurophysiological measures with a shift brain plasticity and cortical inhibition, towards normal levels comparable to healthy control participants.31 As basal ganglia hyperactivation has been implicated in focal hand dystonia, using functional MRI (fMRI) techniques,12 motor, premotor, supplementary motor and basal ganglia activity will be analysed using resting state fMRI pretreatment and post-treatment, in addition to comparison of these findings with asymptomatic controls with a proposed shift to normalisation. Advanced imaging methods, using diffusion tensor imaging (DTI), have demonstrated abnormalities in fibre tracts connecting the primary sensorimotor regions to subcortical structures in focal hand dystonia32 and we aim to assess the effect of Vo-complex thalamotomy on these both these pathways in addition to microstructural changes in the pallidothalamic tract. Similarly, these changes will be compared with healthy control participants at baseline with proposed normalisation of fractional anisotropy.

Objectives

Primary objective
The primary objective of this trial is to establish the efficacy of MRgFUS Vo-complex thalamotomy for the treatment of focal hand dystonia using the Arm Dystonia Disability Scale (ADDs)33 and Writer’s Cramp Rating Scale (WCRS)34 or Tubiana Musicians Dystonia Scale (TMDS),35 as appropriate.

Secondary objectives
1. To establish the adverse event profile of MRgFUS Vo-complex thalamotomy for the treatment of focal hand dystonia.
2. To establish the motor cost associated with MRgFUS Vo-complex thalamotomy by comparing standardised measures of upper limb function preprocedure and postprocedure.
3. To compare baseline measures of neuroplasticity and cortical inhibition in individuals with focal hand dystonia and healthy controls, in addition to comparing
the changes in neuroplasticity and cortical inhibition prethalamotomy and post-thalamotomy.

4. To compare baseline radiological measures using DTI and fMRI in individuals with focal hand dystonia and healthy controls, in addition to comparing the changes in these radiological measures prethalamotomy and post-thalamotomy.

METHODS AND ANALYSIS

Study design
This trial is an open-label clinical trial. Recruitment commenced in November 2022 and is expected to finish in December 2023.

MRgFUS Vo-complex thalamotomy
The MRgFUS intervention will be performed according to the following protocol: on the day of treatment, participants will undergo a head shave and the administration of local anaesthetic blocks prior to application of a stereotactic head frame. An elastic diaphragm is then placed over the scalp and affixed to the ultrasound transducer prior to being filled with cooled, degassed water. Participants are then placed into the 3-Tesla MRI scanner (SIGNA Architect, General Electric) and 3D T1-weighted imaging performed for localisation of the ventro-oral anterior and ventro-oral posterior nuclei. Lesions are created using an Exablate Neuro (InSightec) 650 Hz, 1024-element, phased array ultrasound transducer. Initial operative planning is performed with a series of low power sonications producing mean temperatures of 40°C–45°C to confirm accurate focusing in three orthogonal planes with MR thermography. MR thermometry is used to measure the maximum temperature at target’s central voxel, centring on the target. Following confirmation of accurate targeting, therapeutic sonications are performed by gradually increasing the ultrasound energy and monitoring both the maximal and average temperatures at the target for each sonication. Clinical examinations are performed between each sonication to assess for focal hand dystonia and for the presence of side effects. This protocol is in keeping with the current TGA approved protocol for the use of MRgFUS in tremor treatment.

Eligibility criteria

Inclusion criteria
Participants will be included based on a formal diagnosis of focal task-specific action hand dystonia according to the MDS 2013 classification of dystonia consensus paper.

Individuals may be included if there is mild paroxysmal or persistent dystonia only if the dystonia was noted to be task-specific at onset. Participants must be at least 18 years of age and be willing to provide written consent to participate in the study. Participants must have tried and failed at least one first line therapy for focal hand dystonia, including hand therapy, oral medications and/or botulinum toxin injections. All participants must be ambulant at the time of the procedure.

Exclusion criteria

- History of ischaemic stroke or transient ischaemic event.
- History of stereotactic cerebral surgery.
- Diagnosis of unstable cardiac disease.
- Diagnosis of psychiatric disease.
- Cognitive impairment as indicated by a Mini-Mental State Examination (MMSE) score of <24.
- Coagulopathy or on anticoagulation that cannot be withheld for the intervention.
- Skull density ratio of <0.3.
- Absolute contraindication to MRI (eg, non-MR compatible pacemaker/defibrillator or other non-MR compatible device in situ).
- Relative contraindication to MRI (such as severe claustrophobia) will be assessed on a case-by-case basis.
- Unwilling to provide written informed consent.

Rationale for study design
This trial is an open-label clinical trial study design with baseline comparison to age and gender matched controls. This design allows for comparison of differences in kinematic, neurophysiological and imaging based parameters that exist individuals with focal hand dystonia compared with ‘healthy’ controls. Further, prospective follow-up and data analysis provide insights into treatment efficacy, safety and examines the network changes underpinning improvement in focal hand dystonia.

Treatment groups
This study will be performed on a single cohort of patients with the aim of recruiting ten participants to the open-label trial. All participants have, or will be, included in the trial based on a formal diagnosis of focal hand dystonia made by a consultant neurologist with specialist expertise in movement disorders. The clinical syndromes of focal hand dystonia will include, but not be limited to, writer’s cramp and musician dystonia, with other forms of task specific focal hand dystonia eligible for screening towards study inclusion.

Following enrolment, all intervention participants undergo baseline assessment followed by MRgFUS Vo-complex thalamotomy the following day. A group of 10 age-matched control subjects will be recruited to undergo baseline MRI, neurophysiological and clinical/kinematic testing to establish whether differences exist at baseline. These individuals will not undergo the remaining assessments or follow-up. The control participants will sign a consent form and be made aware that they will not undergo any procedure or follow-up thereafter.

Data collection methods
Data collection will occur at baseline and at 1, 3, 6 and 12 months postprocedure (figure 1). Baseline demographic data (age, gender, hand dominance) will be recorded as standard. Cognitive screening, using the MMSE and mood disturbance, using the Beck Depression Inventory,
will be obtained in all participants. The remaining data will be collected as outlined in table 1.

**Primary outcome**

For the primary outcome, treatment efficacy will be based on administration of the ADDS, either the TMDS or WCRS depending on the syndromic diagnosis, in addition to the Michigan Hand Outcomes Questionnaire and Short Form-36 scores of quality of life, at 1, 3, 6 and 12 months post-treatment. All testing and scoring will be undertaken by study authors JM and ST.

**Secondary outcomes**

The adverse event profile of MRgFUS Vo-complex thalamotomy will be assessed using standardised recording sheets applied on days 0 and 1 postprocedure, in addition to all study timepoints. To assess the motor cost associated with Vo-complex thalamotomy, measures of grip and pincer strength (measured in kg force) will be undertaken at baseline and all time points in addition to validated measures of upper limb function using the box-and-block test, Purdue 9-hole peg and grooved peg tests.

Measures of neuroplasticity will be derived using transcranial magnetic stimulation and paired associative stimuli to obtain pretreatment and post-treatment measures of long-term potentiation-like plasticity with short-latency afferent facilitation, spatial specificity, short-latency intracortical inhibition and stimulus-response curves. These measures will be compared with control participants, at baseline, to assess for any differences in individuals with focal hand dystonia prior to intervention. Repeated measures will be obtained at regular intervals post-treatment as outlined above to identify the effect of MRgFUS Vo-complex thalamotomy on these measures of plasticity.

Radiological outcomes will be assessed by DTI of the pallidothalamic and surrounding tracts to assess for

**Table 1** Data collection and time points

<table>
<thead>
<tr>
<th>List</th>
<th>Enrolment/baseline</th>
<th>Day 0</th>
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<th>3 months</th>
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<td>Questionnaires (ADDS, MHOQ, TMDS/WCRS, SF-36, MMSE, BDI)</td>
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ADDS, Arm Dystonia Disability Scale; BDI, Beck Depression Inventory; MHOQ, Michigan Hand Outcomes Questionnaire; MMSE, Mini-Mental State Examination; SF-36, Short Form-36; TMDS, Tubiana Musicians Dystonia Scale; WCRS, Writer’s Cramp Rating Scale.
Sample size and statistical analysis
A preliminary sample-size calculation indicates that we will require eight patients to detect a 50% improvement in focal hand dystonia (based on standardised rating scales and a predicted similar outcome to a recent pilot study) at p<0.05 with 0.9 power and an SD of 0.30. Ideally, we aim to recruit at least 10 patients to the trial and possibly more pending patients fulfilling inclusion criteria. The data will be analysed using analysis of variance (ANOVA) with main effect of time at 1, 3, 6 and 12 months follow-up intervals. Post hoc paired t-tests (or Shapiro-Wilk if the data is non-normative) will be used where effects of time are identified by ANOVA. Non-parametric variables will be compared pre and post using Wilcoxon signed-rank test.

Patient and public involvement
Patients and the public were not involved in the design of this trial due to its early clinical phase model. Patients were involved in the trial at the time of recruitment and were consulted for their opinion on methods of assessment of fine motor tasks and task performance. Participants in this trial will be primarily recruited from the neurology and movement disorders clinics of the Department of Neurology at St Vincent’s Hospital Sydney. The study treatment costs are fully funded and no financial cost will be borne by participating patients. Additionally, the trial has been advertised through the Brain Foundation, with specific reference made to the trial during a presentation on dystonia given by author JM for Brain Awareness week 2023. The trial details have been disseminated by both the Dystonia Network of Australia and the Australia Dystonia Support Group to all active members. Further, this trial was featured on the Australian Broadcasting Corporation’s nationally televised programme ‘7:30’ on the 26 June 2023. Once the trial results have been published, participants will be informed by the study authors by a dedicated study newsletter.

DISSEMINATION
All adverse events, both non-serious and serious, are recorded and reported to the HREC by the study investigators. All data will be stored securely for five years, in accordance with the National Statement on Ethics Conduct, as well as NSW (New South Wales) health data storage requirements under the Health Records and Information Privacy Act, and will be destroyed thereafter. Paper records will be disposed of in confidential recycling bins and electronic files permanently deleted.

Given its clinical importance, the study will be presented for publication in high-quality neurology journals with a movement disorder subinterest. The strength of this study is in its relative novelty (only the second group worldwide to commence MRgFUS Vo-complex thalamotomy for focal hand dystonia) and the depth of analysis of motor outcomes to fully assess positive and negative impacts of treatment, as well as the corresponding structural and neurophysiological changes underlying those outcomes. Further, all results will be published with the ANZCTR at anzctr.org.au.

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