**Introduction**

Dysphagia is a major complication following acute neurological and more chronic neurodegenerative disorders, resulting in increased risk of death, pneumonia, dehydration, and malnutrition. Rehabilitation-based interventions have been proposed, and therapeutic protocols have been formed with the scope of targeting cortical and subcortical brain areas that are recruited during the highly coordinated sensorimotor activity of swallowing. These paradigms are based on knowledge from in vivo animal and human studies with electrophysiological methods, transcranial magnetic stimulation (TMS), and functional magnetic resonance imaging.

Recently, paired associative stimulation (PAS) paradigms have been developed and investigated on both neurophysiological and behavioral measures of swallowing performance. In this paradigm, pairing a peripheral (pharyngeal electrical stimulation, PES) with a central cortical (TMS) stimulus to the pharyngeal motor cortex (MI) resulted in an increase of cortical excitability of corticomotor projections to the pharyngeal muscles, followed by beneficial behavioral changes. Moreover, following a single application of PAS to the unaffected hemisphere of chronic stroke patients, cortical excitability was increased bilaterally. This was accompanied also by significant functional changes in swallowing physiology and a reduction in the incidence of penetration and/or aspiration of material into the trachea.

However, the effective application of PAS to a dysphagic stroke population may be confounded by several other parameters such as the heterogeneity in responses because of different lesion loci and volumes as well as genetic
factors proposed to be influential in the responsiveness to neurostimulation.\textsuperscript{7} Other potential reported parameters for differences in responses have been the inherent intrinsic neuronal activity,\textsuperscript{8,9} time of day for the intervention delivery,\textsuperscript{10} attentional state,\textsuperscript{11} and cortical thickness in primary sensorimotor cortex.\textsuperscript{12}

This variability in responsiveness has been observed in the literature for the limbs of both in healthy participants\textsuperscript{13} and in stroke patients.\textsuperscript{14} In the latter patient population study\textsuperscript{14} with 9 hemiparetic stroke patients, functional improvements in motor performance were found, while the group’s variability in responses did not allow for significant changes in neurophysiological measurements. Although this has not been the case for our studies in dysphagic stroke patients,\textsuperscript{5} further investigation into other parameters of this neurostimulation paradigm is imperative, since such knowledge will guide us to address the robustness of PAS for swallowing rehabilitation. In point of fact, one controlled study in 5 healthy subjects showed that lithium (a mainstream medication for bipolar disorders) reversed the cortical excitability of “nonresponders” after excitatory PAS into that similar to “responders.”\textsuperscript{15}

Therefore, we investigated the variability in the excitatory responses following the application of the neurorehabilitation paradigm, PAS, in healthy volunteers. Following this initial study, we investigated whether repetitive dosing of PAS could address the variability in responses at the neurophysiological level.

**Participants and Methods**

No major illnesses were reported by the 18 healthy participants (4 men; age, $39 \pm 3$ years [mean $\pm$ SEM]; 16 right-handed). Written informed consent was obtained from all participants before the experiments. General practitioners were informed of the participants’ consent prior to the studies. Exclusion criteria included a history of epilepsy; cardiac pacemaker; previous brain or ear, nose, and throat surgery; any history of swallowing problems; significant medical disorders; pregnancy; metal in the head or eyes; or use of medication that acts on the central nervous system. Research protocols were approved by Salford and Trafford Research Ethics Committee, and experiments were undertaken in the clinical laboratories of the Inflammation Sciences Research Group at Salford Royal NHS, UK, in accordance with the Code of Ethics of the World Medical Association (Declaration of Helsinki).

**Experimental Procedures**

**Transcranial Magnetic Stimulation**

Focal TMS was performed using a flat figure-of-8-shaped magnetic coil (outer diameter 70 mm) connected with a Magstim BiStim\textsuperscript{2} magnetic stimulator (Magstim Co, Whitland, Wales, UK), which produced maximal output of 2.2 T.

**Pharyngeal and Thenar Electromyographic Measurements**

Pharyngeal electromyographic measurements after single TMS pulses, termed pharyngeal motor evoked potentials (PMEPs), were recorded through a 3.2-mm diameter intraluminal catheter (Gaeltec Ltd, Isle of Skye, Scotland) with a built-in pair of bipolar platinum ring electrodes, which was inserted either nasally (15-17 cm to pair electromyographic electrodes from the nasal flare) or orally (13-15 cm) depending on subject’s preference. This allowed the recording of PMEPs at the mid-pharyngeal level (middle pharyngeal constrictors).

As a control (unilaterally innervated) system, thenar motor evoked potentials (TMEPs) from the abductor pollicis brevis muscle were also recorded from MI (see supplementary material).

**Paired Associative Stimulation**

Paired associative stimulation was delivered by pairing a pharyngeal electrical stimulus (0.2-ms pulse) with a single TMS pulse on the pharyngeal MI at the intensity of resting motor threshold (rMT) plus 20% of magnetic stimulator output (MSO). The 2 paired pulses were delivered in a controlled manner through Signal software (v4.1, Cambridge Electronic Design, Cambridge, UK), with an interstimulus interval of 100 milliseconds, based on previous investigations.\textsuperscript{5,6} The intraluminal catheter used for PMEPs was connected to a constant current generator (model DS7; Digitimer, Herts, UK) to deliver pharyngeal electrical stimulation (PES). The paired pulses were delivered every 20 seconds for a total of 10 minutes, giving 30 paired pulses in total. For the sham intervention, the coil was held tangentially to the skull at a 90° angle to sagittal plane, and no PES was delivered through the catheter in situ (see supplementary material).

**Experimental Protocols**

**Protocol I: Real and Sham PAS on Pharyngeal Corticobulbar Projections**

The participants were initially asked to attend the laboratory on 2 occasions. At each attendance, volunteers sat comfortably in a reclining chair with the catheter in situ. The cranial vertex was identified\textsuperscript{16} and marked on the scalp.

The cortical sites for pharyngeal response, characterized as the sites evoking the largest pharyngeal responses in each hemisphere, were identified with mapping procedures using
single TMS pulses delivered over multiple points at 80% MSO intensity, as previously described. The “stronger” pharyngeal projection was defined as the hemispheric site with the lowest rMT to evoke PMEPs, whereas the site with the highest rMT was termed “weaker” pharyngeal projection (see supplementary material).

To assess the effects of real and sham PAS, all participants were studied at least 1 week apart and received 10 minutes of PAS (PAS$_{10\text{min}}$) or sham (PAS$_{\text{Sham}}$) in a randomized manner using block randomization (StatsDirect Ltd, Cheshire, UK). Measurements of cortical excitability for each hemispheric site (10 pulses at rMT + 20% MSO at stronger pharyngeal, weaker pharyngeal, and thenar representation) were made at baseline and at each of the postintervention follow-up time points (immediately, 30, 60, and 90 minutes) on each visit. During these periods, participants were advised to withhold from any swallowing, coughing, talking, or moving their hands or arms. The lead researcher performed the recordings and the analysis but was blinded to the interventions, delivered by a separate researcher who was blinded to the analysis. Participants’ data were kept unidentifiable.

**Protocol 2: Effects of Repeated PAS Over Pharyngeal MI in Responders and Nonresponders**

Following the completion of protocol 1, 12 participants from that protocol were recruited and stratified into 2 groups based on their responses to stimulation of the “stronger” projection (area under the curve [AUC] analysis for PAS$_{10\text{min}}$). Six subjects whose responses were at ≥75th percentile of AUC results after PAS$_{10\text{min}}$ to the “stronger” pharyngeal projection were termed “responders,” whereas the 6 subjects with ≤25th percentile AUC were termed “nonresponders.” The procedures for recording PMEPs and TMEPs, randomization and blinding were identical to protocol 1. Both groups of “responders” and “nonresponders” underwent

(a) double dose of PAS$_{10\text{min}}$ (1 hour intertreatment interval) (Repeat PAS$_{10\text{min}}$),
(b) single dose of PAS$_{10\text{min}}$ followed by PAS$_{\text{Sham}}$ (PAS$_{10\text{min}}$ + PAS$_{\text{Sham}}$), and
(c) PAS$_{\text{Sham}}$ followed by PAS$_{\text{Sham}}$ (PAS$_{\text{Sham}}$ + PAS$_{\text{Sham}}$)

on 3 occasions. Cortical excitability was assessed for up to 60 minutes after the second dose of PAS and then immediately, 30, 60, and 90 minutes after second PAS$_{10\text{min}}$ or PAS$_{\text{Sham}}$.

**Data Analysis of Neurophysiological Measurements**

Peak-to-peak amplitudes of MEPs evoked by TMS were used as a measure of cortical excitability. The individual MEPs were reviewed with Signal Software (CED, Cambridge, UK), MEPs averages were calculated for each time point, and an average trace was created (for response latencies measurements). Baseline MEP data and response latencies for all interventions were compared with nonparametric tests (Friedman test and Wilcoxon signed rank test). Data were normalized to baseline and are shown as percentage change from baseline to minimize interindividual variability. Interindividual factors such as age and sex were therefore equalized. Changes in excitability over time were compared (excluding baseline) using a generalized linear model repeated-measures analysis of variance (rmANOVA; SPSS 16.0). In addition, AUC from percentage change analysis was employed to show the integrated magnitude of the responses of the participants, thus eliminating time-dependency effects. A $P < .05$ was taken as a measure of statistical significance. All data are presented as group mean ± SEM, unless stated otherwise.

**Results**

**Protocol 1: Real and Sham PAS on Pharyngeal Corticobulbar Projections**

PMEPs were recorded in all subjects without any adverse incidents. Larger pharyngeal responses were found from the right hemisphere in 5 participants, whereas the remaining subjects had larger responses from the left hemisphere. The optimal site for stimulation anterior to the vertex was located at 4.6 ± 0.2 cm for the right and 4.9 ± 0.2 cm for the left hemisphere and lateral to midline was 3.8 ± 0.6 cm (right) and 3.7 ± 0.6 cm (left). The mean value of pharyngeal rMT for the “stronger” pharyngeal projection, where PAS was applied, was 67% ± 3% MSO. PES as part of PAS was delivered at 16.6 ± 3.5 mA.

**Baseline TMS response amplitudes and latencies.** Baseline cortical excitability for the 2 different studies remained stable for pharyngeal and thenar projections (see supplementary material).

**Changes in cortical excitability.** A 3-way rmANOVA on percentage change after PAS$_{10\text{min}}$ and PAS$_{\text{Sham}}$ with factors of Intervention, Time, and Site (strong pharyngeal, weak pharyngeal, thenar projection) revealed a significant Intervention × Time × Site interaction ($F_{1,17} = 6.83; P = .018$) and was further analyzed below.

**Changes in PMEP-strong.** A 2-way rmANOVA on the percentage change with the factors: Intervention (PAS$_{10\text{min}}$, PAS$_{\text{Sham}}$) and Time revealed significant Time × Intervention interaction ($F_{1,17} = 6.37; P = .022$) and a significant effect of intervention for PAS$_{10\text{min}}$ against PAS$_{\text{Sham}}$ ($F_{1,17} = 16.22; P = .001$). Compared with PAS$_{\text{Sham}}$, PAS$_{10\text{min}}$ increased cortical excitability, (maximum of 62% ± 23%, 60 minutes). PMEP amplitudes increased significantly immediately ($P = .012$; 95% confidence interval [CI] −87.02 to −12.3) and at 30 minutes ($P = .01; 95\% CI = −52.8 to −7.04$) after PAS$_{10\text{min}}$ compared with baseline. Cortical excitability was still significantly increased up to 51% ± 20% ($P = .04; 95\% CI = −72 to −0.06$) at 90 minutes.
Changes in PMEP-strong. For the “weaker” (nonstimulated) pharyngeal projection, a significant Time × Intervention interaction was observed ($F_{1,11} = 6.6; P = .02$), but there were no significant effects of Intervention or Time.

Changes in TMEP (control). TMEP response amplitudes and latencies following PAS$_{10min}$ and PAS$_{Sham}$ were unaffected (see supplementary material).

AUCs—strong and weak. Nonparametric statistical test (Friedman test) on AUCs of percentage change after PAS$_{10min}$ and PAS$_{Sham}$ showed significant differences in distribution ($P < .001; \chi^2 = 19.6$). Wilcoxon tests performed on AUC after PAS$_{10min}$ and PAS$_{Sham}$ showed significant difference only for the “stronger” pharyngeal projection compared with PAS$_{Sham}$ ($z = -3.33; P = .001$), verifying the aforementioned results (see Table 1 and Supplementary material). The subjects were then stratified to “responders” ($\geq$75th percentile) and “nonresponders” ($\leq$25th percentile).

Protocol 2: Effects of Repeated PAS Over Pharyngeal MI in Responders and Nonresponders

Twelve volunteers (11 women; age, 43 ± 8 years, mean ± SEM) were invited to participate in protocol 2, and Figure 1 shows their responses for PAS$_{10min}$ on the “stronger” pharyngeal projection. As with protocol 1, all measurements in protocol 2 were recorded with no adverse incidents (see supplementary material).

Cortical excitability changes for “responders” and “nonresponders”. Baseline TMS response amplitudes were similar and latencies remained unaffected across the 3 arms for all sites (see supplementary material).

Changes in cortical excitability. A 3-way rmANOVA with factors Intervention (Repeat PAS$_{10min}$ × PAS$_{Sham}$, PAS$_{Sham}$ × PAS$_{Sham}$, PAS$_{Sham}$ + PAS$_{Sham}$), Time (immediately, 30, 60, and 70 minutes; 90, 120, and 150 minutes) and Site (strong, weak, and thenar projection) showed a significant interaction of Intervention × Time × Site interaction ($F_{1,11} = 8.60; P = .016$). Three separate 2-ways rmANOVAs were performed for each hemispheric Site with factors Intervention and Time.

Changes in PMEP-strong. Figure 2A shows the changes in cortical excitability of the stimulated—“stronger” pharyngeal projection in all subjects for the 3 different studies. There was a significant Time × Intervention interaction ($F_{1,11} = 20.4; P = .001$). Moreover, there was a significant difference between the interventions of repeated PAS$_{10min}$ versus PAS$_{Sham}$ ($F_{1,11} = 19.5; P = .001$) and a significant difference between repeated PAS$_{10min}$ versus PAS$_{Sham}$ ($F_{1,11} = 8.7; P = .013$). The effect of the application of single PAS$_{10min}$ was also significantly different compared with PAS$_{Sham}$ ($F_{1,11} = 9.15; P = .012$).

The maximum increase in group percentage change (up to 95% ± 29%) was observed 60 minutes after repeated PAS$_{10min}$ whereas after the single application of PAS$_{10min}$, maximum percentage change reached 25% ± 18% at 60 minutes (Figure 2A).

Changes in PMEP-weak. There was a significant interaction of Time × Intervention for the “weaker” (nonstimulated) pharyngeal site ($F_{1,11} = 6.2; P = .029$). However, only the effects of repeated PAS$_{10min}$ were significantly different compared with PAS$_{Sham}$ ($F_{1,11} = 6.6; P = .025$) with excitability

| Table 1. Group Changes in Cortical Excitability After Real and Sham PAS. |
|-------------------------|----------|------------|----------|----------|----------|----------|----------|
|                         | 25th Percentile | Median     | 75th Percentile | 25th Percentile | Median     | 75th Percentile |
| **Strong Projection**   |          |            |            |          |            |            |
| PAS$_{10min}$           | 15.67    | 148.9      | 235.9      | -68.9    | 0.89      | 85.5      |
| PAS$_{Sham}$            | -104.2   | -55.9      | -27.0      | -109.8   | -40.5     | 8.22      |
| **Weak Projection**     |          |            |            |          |            |            |

Abbreviations: PAS, paired associative stimulation; PMEP, pharyngeal motor evoked potential.

aGroup mean area under the curve (calculated by the percentage change in PMEPs’ amplitude against time) analysis of “stronger” and “weaker” pharyngeal projection after PAS$_{10min}$ and PAS$_{Sham}$. There was a significant difference between the change in cortical excitability of the “stronger” projection following real PAS compared with sham ($z = -3.33, P = .001$).
maximally increasing to 34% ± 22% at 90 minutes, whereas after the single PAS, this was 2.6% ± 9% at 30 minutes after the initial PAS (Figure 2B).

Changes in TMEP. Application of repeated and single PAS did not change thenar muscle excitability as compared with sham (see supplementary material).

Responders versus nonresponders. The percentage changes for “responders” and “nonresponders” after repeated and single PAS are shown in Figure 3.

Two different AUC analyses were performed and compared with nonparametric tests. We first included each group’s responses for all time points up to the end point (method A). We then calculated the effects of each application (PAS or PAS_sham) up to 60 minutes following baseline and up to 90 minutes following the second application of PAS or PAS_sham (method B).

Method A. Friedman χ² was 39.7 and gave P values of <.001 (stronger projection) and 10.3 with P = .006 (weaker projection), suggesting that the distributions were different for both pharyngeal projections. Nonparametric tests were then performed to capture the differences between “responders” and “nonresponders.” Table 2 presents the different responses of the group of “responders” and “nonresponders” for both projections across all interventions.

Method B. Repeated application of PAS further increased the excitability for responders compared with the initial application for the “stronger” projection (z = −2.2; P = .02). The effects of both initial and repeat PAS were significantly different compared with PAS_sham both for the stronger and weaker projections for “responders” (both: z = −2.2; P = .02). Importantly, responders’ AUCs after repeated PAS were also increased compared with single PAS for the same period (z = 1.94; P = .046). As expected, the effects of single PAS were significantly different compared with sham for both the initial 60 minutes and for up to 150 minutes for the “responders,” indicative that in “responders” a single application of PAS may induce long-term effects.

Repeated PAS also resulted in an increase to the stronger pharyngeal projection in “nonresponders” (z = −2.2; P = .02), which was significantly different when compared with single PAS and PAS_sham (z = −2.2; P = .02). There was no difference between the effects of the single active PAS and

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**Figure 2.** Group mean percentage change in PMEPs amplitude, on the ‘stronger’ (stimulated) (A) and ‘weaker’ (B) pharyngeal projection following different PAS doses and sham stimulation. Increase in amplitude in the ‘stronger’ pharyngeal projection was observed following both repeated PAS (♦) (P=.001) and single PAS (■) for the initial period after the first application up to 60 minutes (P=.012), compared to sham PAS (●). For the ‘weaker’ pharyngeal projection (B) only repeated PAS resulted in significant increase in cortical excitability (P=.025). Abbreviations: PMEP, pharyngeal motor evoked potential; PAS, paired associative stimulation.
the effects after PAS$_{\text{Sham}}$ or the effects of the first period of stimulation in the double dose PAS arm for the “nonresponders,” in keeping with previous results for the reduced effects following single PAS to “nonresponders” in protocol 1.

**Discussion**

The effects of PAS$_{10\text{min}}$ on the “stronger” pharyngeal projection corroborate the results of our previously published data.
and show that PAS has the potential to excite the swallowing neural network. Most important, this study set out to examine the effects of repeated PAS $_{\text{10min}}$ in 2 groups of subjects in whom PAS was either excitatory or ineffective and to investigate whether PAS repetition could further modulate MI excitability. Our observation that additional doses of PAS have the potential to convert “nonresponders” to “responders” is of interest and merits further discussion.

**Dose Effects of PAS on Bilateral Pharyngeal MI**

Repeated PAS $_{\text{10min}}$ over the “stronger” pharyngeal projection in both “responders” and “nonresponders” induced facilitation in both stimulated and unstimulated hemispheres, with cortical excitability in the stimulated MI being significantly increased after the second application. The magnitude of these facilitatory effects is surprising, since the group consisted of equal numbers of “responders” and “nonresponders.” Separate analysis for the effects of repeated PAS to “responders” and “nonresponders” individually (controlled with single and sham PAS) indicated that the effects were mainly because of the second PAS.

Previous work on limb muscles in healthy volunteers, stroke patients, and animal models have shown that repeated PAS once per day for 5 days or even longer (ie, in stroke patients) enhanced neurophysiological properties of the corticomotor system, as measured by MEP amplitude, accompanied with behavioral benefits. However, results from our studies are not directly comparable, since the repeat PAS protocol was applied within a shorter epoch to the initial intervention.

Notwithstanding, the findings from our current study differ from those by Müller et al. These authors found that when cortical excitability was conditioned with a PAS paradigm that enhances long-term potentiation (LTP), then the application of a second LTP-like PAS intervention resulted in cortical depression. The results from that study fall within the well-described theory of Bienenstock–Cooper–Munro, which tries to elucidate the way that neuronal systems reach homeostasis and balance inhibitory and facilitatory interactions over a period of time, originally observed in visual cortical neurons. In contrast, our data have shown that the effect of the first PAS $_{\text{10min}}$ application resulting in LTP-like plasticity in pharyngeal motor cortex was further enhanced after a second facilitatory PAS $_{\text{10min}}$ application. This finding requires further consideration.

There are likely to be a number of explanations for the difference in the results in the swallowing model. First, the existence of the “ceiling effect,” the extent to which cortical excitability can be further increased, has not been examined for the swallowing motor system. Second, the effect of “saturation” of the cortical capacity for synaptic efficacy and LTP has also not been investigated in detail for swallowing. However, previous PAS studies showed that 30 minutes of stimulation did not produce significant changes compared with shorter durations. The inter-PAS interval is also an important parameter to consider for the modulatory effects of PAS to MI. In the study by Müller et al., the inter-PAS interval was 30 minutes, shorter than the 60-minute inter-PAS interval in our protocol. Furthermore, we have previously shown that the effects of single PAS targeting pharyngeal MI can last up to 90 minutes. In this context, evidence from the use of transcranial direct current stimulation in the limb MI has shown that when the repeated application falls within the excitatory window of the initial input, the after effects are increased. In addition, the interval between the pairs of peripheral and cortical stimulation is critically important: Literature by others has suggested that with different intervals between pairs, different mechanisms contribute to the effects of PAS.

Moreover, at this stage of research, it is still unclear as to whether the changes in cortical excitability are because of changes in the efficacy of the synaptic connections or changes in neuronal excitability. Cortical and subcortical brain areas are activated in an interconnected network during swallowing. Whether the change in cortical excitability following the first application of PAS would spread to connected brain regions of the swallowing network resulting in a further increase after the application of repeat PAS is uncertain. Further studies with neuroimaging techniques may help determine this and would be of importance for the applications of neurorehabilitation to the corticobulbar network for swallowing.

More recently, it has been shown that PAS effects in limb MI can be remotely influenced by cerebellar stimulation. Modulation of cerebellar activity using transcranial direct current stimulation was able to abolish the excitatory effects of PAS in the motor cortex. These findings suggest that combining neurostimulation inputs is both modality and region specific, which supports the contention that pharyngeal motor cortex neurostimulation might behave differently in other regions, when PAS is applied.

Nonetheless, our study also raises the possibility that delivering initial PAS as a form of conditioning changed the threshold for synapses to engage in “nonresponders” (producing an “imbalance” in activity), and the repeat PAS has enabled these “activated” synapses to be strengthened more easily. However, in vivo studies to validate this assumption are difficult to perform, given that the measurement of excitability is not directly equal to synaptic activity. Nevertheless, such suggestion could hold considerable value for the rehabilitation of swallowing disorders, if we take into consideration that LTP induced by targeted PAS, such as in our study, has similarities with LTP resulting from motor training and learning, the latter being important in the case of dysphagia rehabilitation.

In conclusion, we report evidence that subjects who do not respond to an initial application of excitatory stimulation (PAS $_{\text{10min}}$) can show an increase in MEP responses after...
a repeated excitatory PAS\_{\text{10min}}; these effects being larger than when compared with a single application. This has implication for PAS application to dysphagic stroke patients who may not respond to single doses of stimulation and provides the example for other neuromodulatory interventions under investigation for customized approaches when applied to the swallowing neural network. Future utilization of the repeated approach in stroke patients with dysphagia and neuroimaging studies seem warranted, since PAS appears to hold promise as a powerful neuromodulation paradigm for dysphagia rehabilitation after stroke. Double PAS could therefore drive cortical plasticity during the critical period of plasticity in the weeks following a stroke and may substantially enhance traditional therapy.\textsuperscript{21}

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