Review

Behavioural Interventions in People with Oropharyngeal Dysphagia: A Systematic Review and Meta-Analysis of Randomised Clinical Trials

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Abstract: Objective: To determine the effects of behavioural interventions in people with oropharyngeal dysphagia. Methods: Systematic literature searches were conducted to retrieve randomized controlled trials in four different databases (CINAHL, Embase, PsycINFO, and PubMed). The methodological quality of eligible articles was assessed using the Revised Cochrane risk-of-bias tool for randomised trials (RoB 2), after which meta-analyses were performed using a random-effects model. Results: A total of 37 studies were included. Overall, a significant, large pre-post interventions effect size was found. To compare different types of interventions, all behavioural interventions and conventional dysphagia treatment comparison groups were categorised into compensatory, rehabilitative, and combined compensatory and rehabilitative interventions. Overall, significant treatment effects were identified favouring behavioural interventions. In particular, large effect sizes were found when comparing rehabilitative interventions with no dysphagia treatment, and combined interventions with compensatory conventional dysphagia treatment. When comparing selected interventions versus conventional dysphagia treatment, significant, large effect sizes were found in favour of Shaker exercise, chin tuck against resistance exercise, and expiratory muscle strength training. Conclusions: Behavioural interventions show promising effects in people with oropharyngeal dysphagia. However, due to high heterogeneity between studies, generalisations of meta-analyses need to be interpreted with care.

Keywords: deglutition; swallowing disorders; RCT; intervention; compensation; rehabilitation
1. Introduction

Swallowing disorders, or oropharyngeal dysphagia (OD), can be the result of many underlying conditions such as stroke, progressive neurological diseases, and acquired brain injury. They may also be the consequence of treatment side effects; for example, radiation or surgical interventions in patients with head and neck oncological disorders. Prevalence of OD in the general population ranges from 2.3 to 16% [1]. However, depending on underlying disease severity and outcome measures used (e.g., instrumental assessment, screening or patient self-report) [2], prevalence estimates can be as high as 80% in stroke and Parkinson’s disease patients, up to 30% in traumatic brain injury patients, and over 90% in patients with community-acquired pneumonia [3]. Also, pooled prevalence estimates for swallowing problems in people with cerebral palsy determined by meta-analyses are as high as 50.4% [4].

OD may have severe effects on a person’s health as dysphagia can lead to dehydration, malnutrition, and aspiration pneumonia. OD also has a high disease burden and poses a major societal challenge, which is associated with significant psychological and social burden, resulting in reduced quality-of-life for both patients and caregivers [5].

The treatment of OD may include surgical, pharmacological and behavioural interventions. Behavioural interventions include: bolus modification and management (e.g., adjusting the viscosity, volume, temperature and/or acidity of food and drinks), motor behavioural techniques or oromotor exercises, general body and head postural adjustments, swallowing manoeuvres (e.g., manoeuvres to improve food propulsion into the pharynx and airway protection), and sensory and neurophysiologic stimulation (e.g., neuromuscular electrical stimulation [NMES]) [6].

An increasing number of reviews have been published over the last two decades on the treatment effects of behavioural interventions in people with OD. However, only one systematic review [7] summarised the effects of swallowing therapy as applied by speech and language therapists without restrictions on subject populations or study designs. Furthermore, while most reviews have focussed on selected types of interventions and patient populations, very few reviews use criteria related to study designs (e.g., [8,9] solely including randomised controlled trials [RCTs], ranked as the highest level of evidence [10]).

This systematic review aimed to determine the effects of behavioural interventions in people with OD based on the highest level of evidence (RCTs) only. Behavioural interventions comprised any intervention by a dysphagia expert, excluding surgical and pharmacological interventions. Clinicians being referred to as dysphagia experts include speech therapists, occupational therapists, or physiotherapists, but may incorporate other disciplines depending on national healthcare and education systems. Finally, neurostimulation techniques were considered out of scope of this current review.

2. Methods

The methodology and reporting of this systematic review were based on the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) statement and checklist. The PRISMA 2020 statement and checklist (Supplementary Tables S1 and S2) aim to enhance the essential and transparent reporting of systematic reviews [11,12]. The protocol for this review was registered at PROSPERO, the international prospective register of systematic reviews (registration number: CRD42020179842).

2.1. Information Sources

To identify studies, literature searches were conducted on 6 March 2021, across these four databases: CINAHL, Embase, PsycINFO, and PubMed. Publications dates ranged from 1937–2021, 1902–2021, 1887–2021, and late 1700s–2021, respectively. Additional searches included checking the reference lists of eligible articles.
2.2. Search Strategies

Electronic search strategies were performed in all four databases using subheadings (e.g., MeSH and Thesaurus terms) and free text terms. Two strings of terms were combined: (1) dysphagia and (2) randomised controlled trial. The full electronic search strategies are reported in Table 1.

### Table 1. Search strategies.

| Database and Search Terms                                                                 | Number of Records |
|------------------------------------------------------------------------------------------|-------------------|
| Cinahl: ((MH “Deglutition”) OR (MH “Deglutition Disorders”)) AND (MH “Randomized Controlled Trials”) | 239               |
| Embase: (swallowing/OR dysphagia/) AND (randomization/or randomized controlled trial/OR “randomized controlled trial (topic)”)/OR controlled clinical trial/ | 4550              |
| PsycINFO: (swallowing/OR dysphagia/) AND (RCT OR (Randomised AND Controlled AND Trial) OR (Randomised AND Clinical AND Trial) OR (Randomised AND Clinical AND Trial) OR (Controlled AND Clinical AND Trial)).af. | 231               |
| PubMed: (“Deglutition”[Mesh] OR “Deglutition Disorders”[Mesh]) AND (“Randomized Controlled Trial” [Publication Type] OR “Randomized Controlled Trials as Topic”[Mesh] OR “Controlled Clinical Trial” [Publication Type] OR “Pragmatic Clinical Trials as Topic”[Mesh]) | 3039              |

2.3. Inclusion and Exclusion Criteria

The following criteria for inclusion were applied: (1) participants had a diagnosis of OD; (2) behavioural interventions were aimed at reducing swallowing or feeding problems; (3) studies included a comparison group; (4) participants were randomly assigned to one of the study arms or groups; (5) studies were published in English.

Studies focussing on drooling, self-feeding, gastro-oesophageal reflux or oesophageal dysphagia (e.g., dysphagia resulting from oesophageal carcinoma or esophagitis) were excluded. Further excluded studies were those describing drug-induced swallowing problems, temporary swallowing problems caused by oedema post-surgery (e.g., anterior cervical discectomy), or swallowing problems associated with adverse effects of interventions such as inflammation and oedema resulting from recent radiotherapy (≤three months after intervention) or thyroidectomy. Studies reporting solely on feeding tube removal after intervention that did not provide data on swallowing or feeding problems, were also excluded. Studies on behavioural eating problems including bulimia, anorexia, and picky eaters, were out of scope of this review. Finally, only original research was included, thus excluding, for example, conference abstracts, doctoral theses and reviews.

2.4. Systematic Review

**Methodological Quality and Risk of Bias.** The Revised Cochrane risk-of-bias tool for randomised trials (RoB 2) [13] was used to assess the methodological quality of the included studies. The RoB 2 tool provides a framework for evaluating the risk of bias in the findings of any type of randomised trial. The tool is structured along five domains through which bias might be introduced into the study results: (1) the randomisation process; (2) deviations from intended interventions; (3) missing outcome data; (4) measurement of the outcome; (5) selection of the reported result.

**Data Collection Process.** A data extraction form was created to extract data from the included studies under the following categories: methodological quality, participant diagnosis, inclusion criteria, sample size, age, gender, intervention goal, intervention agent/delivery/dosage, intervention condition, outcome measures and treatment outcome.
Data, Items and Synthesis of Results. Two independent raters reviewed all titles and abstracts, then original articles, for eligibility. Inclusion of studies was based on consensus between raters. To ensure rating accuracy, two group sessions were held to discuss ratings of one hundred randomly selected records to achieve consensus before rating the remaining abstracts. Where consensus could not be reached between the first two raters, a third party was consulted for resolution. Methodological quality assessment was also rated by two independent researchers, after which consensus was reached with involvement of a third reviewer, when necessary. No evident bias in article selection or methodological study quality rating was present as none of the reviewers had formal or informal affiliations with any of the authors of the included studies. At this stage reviewers did not exclude studies based on type of intervention (e.g., behavioural intervention, neurostimulation).

During data collection, data points across all studies were extracted using comprehensive data extraction forms. Risk of bias was assessed per individual study using RoB 2 [13]. The main summary measures for assessing treatment outcome were effect sizes and significance of findings.

2.5. Meta-Analysis

Data was extracted from relevant studies to compare the effect sizes for the following: (1) pre-post outcome measures of OD and (2) mean difference in outcome measures from pre to post between different types of behavioural interventions. All interventions were categorised into compensatory (e.g., body and postural adjustments, or bolus modification), rehabilitative (e.g., oromotor exercises or Shaker exercise), combined compensatory and rehabilitative interventions, and no dysphagia intervention. Only studies using instrumental assessment (videofluoroscopic swallow study [VFSS] or fiberoptic endoscopic evaluation of swallowing [FEES]) to confirm OD were included. Outcome measures based on visuoperceptual evaluation of instrumental assessment and clinical non-instrumental assessments, were eligible for inclusion in meta-analyses. However, if both types of data were available, instrumental assessment was preferred over non-instrumental assessment outcome data. Oral intake measures, screening tools and patient self-report measures were excluded from meta-analyses. Measures other than the authors’ primary outcomes may have been selected if these measures helped to reduce heterogeneity between studies.

To compare effect sizes, group means, standard deviations, and sample sizes for pre- and post-measurements were entered into Comprehensive Meta-Analysis Version 3.3.070 [14]. If only non-parametric data were available (i.e., medians, interquartile ranges), then data were converted into parametric data for meta-analyses. Participants in studies of multiple intervention groups were analysed separately. Where studies used the same participants, only one study was included in the meta-analysis. If studies provided insufficient data for meta-analyses, authors were contacted by e-mail for additional data.

Effect sizes were calculated in Comprehensive Meta-Analysis using a random-effects model. Due to variations in participant characteristics, intervention approaches, and outcome measurements, studies were unlikely to have similar true effects. Heterogeneity was estimated using the $Q$ statistic to determine the spread of effect sizes about the mean and $I^2$ was used to estimate the ratio of true variance to total variance. $I^2$-values of less than 50%, 50% to 74%, and higher than 75% indicate low, moderate, and high heterogeneity, respectively [15]. Using the Hedges $g$ formula for standardized mean difference with a confidence interval of 95%, effect sizes were calculated and interpreted using Cohen’s $d$ convention: $g \leq 0.2$ as no or negligible effect; $0.2 < g \leq 0.5$ as minor effect; $0.5 < g \leq 0.8$ as moderate effect; and $g > 0.8$ as large effect [16].

Forest plots of effect sizes for OD outcome scores were generated for pre-post behavioural interventions. Due to blended configurations of intervention groupings across studies it was not possible to compare a homogenous behavioural intervention group against a comparison group that did not have a behavioural component. For this reason, only a
A subgroup between group analysis was conducted (and not an overall between group analysis) to explore effect sizes as a function of various moderators. Behavioural interventions (compensatory, rehabilitative, or combined compensatory and rehabilitative interventions) were compared with conventional dysphagia treatment (CDT), or no dysphagia therapy groups. Other subgroup analyses were conducted to compare effect sizes between selected interventions (i.e., Shaker exercise, Chin Tuck Against Resistance exercise [CTAR], and Expiratory Muscle Strength Training [EMST]), medical diagnoses, and outcome measures. Only between-subgroup meta-analyses were conducted using post-intervention data, to account for possible spontaneous recovery during the period of intervention.

Using Comprehensive Data Analysis software, publication bias was assessed following the Begg and Muzumdar’s rank correlation test and the fail-safe N test. The Begg and Muzumdar’s rank correlation test reports the rank correlation between the standardised effect size and the variances of these effects [17]. This statistical procedure produces tau as well as a two tailed $p$ value; values of zero indicate no relationship, whereas deviations away from zero indicate a relationship. High standard error would be associated with larger effect sizes if asymmetry is caused by publication bias. Tau would be positive if larger effects are presented by low values, while tau would be negative if larger effects are represented by high values.

The fail-safe N test calculates how many studies with effect size zero could be added to the meta-analysis before the result lost statistical significance. That is, the number of missing studies that would be required to nullify the effect [18]. If this number is relatively small, then there is cause for concern. However, if this number is large, it can be stated with confidence that the treatment effect, while possibly inflated by the exclusion of some studies, is not nil.

3. Results
3.1. Study Selection

A total of 8059 studies were retrieved across four databases: CINAHL ($n = 239$), Embase ($n = 4550$), PsycINFO ($n = 231$), and PubMed ($n = 3039$). After removal of duplicate titles and abstracts ($n = 1113$), a total of 6946 records remained. After assessing titles and abstracts, 261 original articles were identified. Full-text records were accessed to verify all inclusion criteria. During full-text assessment, articles were divided into different types of interventions, as this systematic review reports on behavioural interventions only. Based on the inclusion criteria, 36 articles were included, after which one study was identified through reference checking of the included articles. Figure 1 presents the flow diagram of the article selection process according to PRISMA.
3.2. Description of Studies

All 37 included studies are described in detail in Tables 2 and 3. Table 2 reports on study characteristics, definitions and methods of diagnosing oropharyngeal dysphagia, and details on participant groups. Information such as medical diagnosis, sample size, age and gender, is provided on all study groups. Table 3 presents intervention goals, intervention components, outcome measures and treatment outcome of each included study.
Table 2. Study characteristics of studies on behavioural interventions for people with oropharyngeal dysphagia.

| Study | Country | OD (Definition/Terminology; Diagnostic Measure/Method) | Sample (N) | Group Descriptive (Mean ± SD) (Age, Gender, Relevant Medical Diagnoses) |
|-------|---------|-------------------------------------------------------|------------|-----------------------------------------------------------------------|
| Ayres, et al. [19] | Brazil | OD: Oropharyngeal dysphagia determined by FEES  
Diagnosis: PD  
Inclusion: PD and oro-phenyngal dysphagia. Exclusion: Presenting language and/or hearing disorders that could complicate the understanding of intervention; diagnosis of dementia, or other neurological illnesses. | n = 32:  
- Experimental group: Chin-down manoeuvre and swallowing orientation (n = 11)  
- Orientation group: Swallowing orientation only (n = 7)  
- Control group: No intervention (n = 14) | Group A/Group B/Group C  
Age: 62 (11.5)/64.5 (5.6)/62.8 (6.2)  
Male: 80%/66.7%/75% |}

| Carnaby, et al. [20] | USA | OD: Diagnosis of swallowing difficulty by speech pathologist, <85 on Hospital’s dysphagia assessment  
Diagnosis: Clinician diagnosed Stroke, WHO definition  
Inclusion: Stroke < 7 days  
Exclusion: NR | n = 306:  
- UC (n = 102)  
- Low intensity (n = 102)  
- High intensity (n = 102) | High intensity/low intensity/LC: mean (SD)  
Age yr: 69.8 (12.5)/72 (12.4)/71.4 (12.7)  
Male: 59%/58%/58%  
Severity Barthel index <15: 85%/79%/83% |}

| Carnaby, et al. [21] | USA | OD: Dysphagia on admission- score < 178 on MASA, no history of swallowing disability, head/neck surgery.  
Diagnosis: Sub-acute stroke confirmed by attending neurologist according to the WHO definition  
Inclusion: Able to adhere to behavioural treatment regimens  
Exclusion: NR | n = 53:  
- MDTP + NMES (NMES; n = 18),  
- MDTP + sham NMES (MDTP; n = 18) [Denoted as ‘Carnaby et al. (2020a)’ in Figure 4.]  
- UC (n = 17) [Denoted as ‘Carnaby et al. (2020b)’ in Figure 4.] | NMES/MDTP/LC: mean (SD)  
Age yr: 62.7 (12.2)/70.6 (11.8)/64.3 (14.7)  
Male: 55%/44%/41%  
Modified Rankin: 4.5 (0.6)/4.46 (0.5)/4.56 (0.5) |}

| Choi, et al. [22] | Korea | OD: Dysphagia after stroke confirmed by VFSS  
Diagnosis: Stroke by clinical history, neurologic examination CT/MRI  
Inclusion: No major cognitive deficit (MMSE ≥ 24), >fair grade on neck muscle testing, symmetric neck posture  
Exclusion: neck pain or neck surgery, poor general condition, severe communication problem, unstable medical condition, presence of a tracheostomy tube | n = 32:  
- Experimental–Shaker exercise (SE) and conventional dysphagia therapy (CDT; n = 16) [Denoted as ‘Choi et al. (2017a)’ in Figure 4.]  
- Control–CDT (n = 16) [Denoted as ‘Choi et al. (2017b)’ in Figure 4.] | Experimental SE + CDT/control (CDT): mean (SD)  
Age yr: 60.8 (10.9)/60.4 (10.5)  
Gender (male/female): 10/69/6  
Time since stroke onset months: 3.4 (1.6)/4.1 (1.0)  
PAS: 4.6 (0.8)/4.9 (0.1)  
FOIS: 3.1 (1.0)/3.2 (0.6) |}

| DePippo, et al. [23] | USA | OD: MBS, BDST, VFSS, speech pathologists determined dysphagia  
Diagnosis: Stroke by clinical history, neurologic examination CT/MRI  
Inclusion: 20–90 yrs, no history of oral or pharyngeal anomaly  
Exclusion: aspirated >50% of all consistencies, | n = 115, allocated to graded therapist treatment levels:  
- Group A (n = 38)  
- Group B (n = 38)  
- Group C (n = 39) | Group A/Group B/Group C  
Age yr: 76/74/73  
Male/Female: 22/16/19/27/12  
Mini-Mental State score: 16 (12)/17 (10)/18 (10)  
Barthel-ADL Mobility: 37 (23)/48 (20)/46 (38)  
Weeks post stroke: 4.6/4.5/4.9 |}

| Eom, et al. [24] | Korea | OD: Dysphagia caused by a stroke, confirmed by VFSS  
Diagnosis: Stroke  
Inclusion: Age ≥ 65, onset duration < 3 months, score ≥ 24 on MMSE. | n = 30:  
- Experimental- resistance expiratory muscle strength training (n = 15)  
- Placebo group (n = 15). | Experimental/Placebo  
Age yr: 69.2 (4.1)/70.2 (3.6)  
Male/Female: 5/8/6/7  
PAS baseline: 5.1 (0.8)/4.9 (0.6) |
| Study | Country | Sample Size | Study Type | Inclusion | Exclusion | OD | Diagnosis | Inclusion | Exclusion | OD | Diagnosis | Inclusion | Exclusion | OD | Diagnosis |
|-------|---------|-------------|------------|-----------|-----------|----|-----------|-----------|-----------|----|-----------|-----------|-----------|----|-----------|
| Gao and Zhang [25] | China | 90 | VFSS evaluation | Chinese diagnosis guidelines for acute ischemic stroke, CT or MRI | Presence of severe orofacial pain, significant malocclusion or facial asymmetry, unstable breathing or pulse, tracheostomy, aphasia or apraxia, inadequate lip closure | n = 30 | Dysphagia confirmed by VFSS score ≥3 in 8-point PAS | Stroke within 1–3 wks. | Cognitive impairment and/or history of previous neurological diseases - associated with dysphagia | n = 32 | Dysphagia confirmed by VFSS score ≥3 in 8-point PAS | Stroke type: | Infarction: 5/4; Haemorrhage: 3/4; Ischemic and ICH = 1/0; Stroke onset (days): 9.3 (5.1)/10.8 (8.7)/11.0 (5.5) | FOIS: 4.3 (0.6)/4.5 (0.5)/4.4 (1.0) | PAS: 5.4 (2.5)/5.2 (2.7)/5.2 (2.2) | Dysphagia risk condition: 32 (47.8)/25 (52.1) | Swallowing rate (mL/s): 4.10/5.31/9.00 | Study/Control: denoted as ‘Goa & Zhang (2017b)’ in Figure 4. | Control/Intervention: Oral neuromuscular training (n = 49) | Age: 85/83 | Male: 29 (43.3)/27 (55.1) | Dysphagia risk condition: 32 (47.8)/25 (52.1) | Care moderate dependence: 27 (40.9)/18 (36.7) | Swallowing rate (mL/s): 4.10/5.31 | Experimental/Control: Experimental, tongue stretching exercises (TSE) (n = 13) | Age: 60.5 (12.5)/62.2 (10.3) | Male: 6/5 | Time since stroke, weeks: 2.2 (9.2)/3.5 (2.6) | Type of stroke (n) Haemorrhage: 7/6 | Type of stroke (n) Infarction: 4/4 | Experimental/Control: Experimental, tongue stretching exercises (TSE) (n = 13) | Age: 60.5 (12.5)/62.2 (10.3) | Male: 6/5 | Time since stroke, weeks: 2.2 (9.2)/3.5 (2.6) | Type of stroke (n) Haemorrhage: 7/6 | Type of stroke (n) Infarction: 4/4 | Experimental/Control: Experimental, tongue stretching exercises (TSE) (n = 13) | Age: 60.5 (12.5)/62.2 (10.3) | Male: 6/5 | Time since stroke, weeks: 2.2 (9.2)/3.5 (2.6) | Type of stroke (n) Haemorrhage: 7/6 | Type of stroke (n) Infarction: 4/4 | Experimental/Control: Experimental, tongue stretching exercises (TSE) (n = 13) | Age: 60.5 (12.5)/62.2 (10.3) | Male: 6/5 | Time since stroke, weeks: 2.2 (9.2)/3.5 (2.6) | Type of stroke (n) Haemorrhage: 7/6 | Type of stroke (n) Infarction: 4/4 | Experimental/Control: Experimental, tongue stretching exercises (TSE) (n = 13) | Age: 60.5 (12.5)/62.2 (10.3) | Male: 6/5 | Time since stroke, weeks: 2.2 (9.2)/3.5 (2.6) | Type of stroke (n) Haemorrhage: 7/6 | Type of stroke (n) Infarction: 4/4 |
| Study/Control | Japan | Poland | Korea | Japan | Poland | Korea | Japan | Korea |
|--------------|-------|--------|-------|-------|--------|-------|-------|-------|
| **Diagnosis:** Stroke | Subacute stroke | Inclusion: Diagnosis of subacute stroke | Exclusion: Previous stroke, pharyngeal structural abnormalities, unable to cooperate | Study-conventional therapy + mechanical inspiration, expiration exercise (n = 18) [Denoted as ‘Jang et al. (2019a)’ in Figure 4.] | Male, n: 10/9 | Study/Control | Male, n: 10/9 | Study/Control | Male, n: 10/9 |
| **Study/Control** | | | | | | | | | |
| **Diagnosis:** Stroke | Stroke type, n Haemorrhage: 8/6 | Days from stroke onset: 20.5 (13.6)/18.4 (12.5) | Male: 12/14 | Paralysis, right side: 15/12 | Total sample | Male: 12/14 | Paralysis, right side: 15/12 | Total sample | Male: 12/14 |
| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
| **OD:** Swallowing dysfunction/dysphagia as determined by VDS and PAS scores on VFSS | Experimental group: NMES + upper cervical spine mobilisation (n = 17) | Control group: NMES and sham mobilization (n = 17) | Male: 11/6, 11/6 | Side of stroke (left/right): 6/11; 7/10 | Haemorrhage/infection: 14/3/12/5 | Weight: 69.11 (11.93); 65.55 (12.66) | K-MMSE (point): 24.53 (2.62)/24.2 (2.91) | K-NHSS (point): 10.41 (3.06)/10.76 (3.75) | | |
| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
| **OD:** Dysphagia defined as a disorder that causes difficulty with chewing and swallowing food | Experimental group: PNF short-flexion neck exercises (n = 13) | Control group: Shaker exercise (n = 13) | Male: 8/5; 7/8 | Side of stroke (right/left): 7/6/7/6 | | | | | |
| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
| **OD:** Dysphagia confirmed by VFSS | Experimental group: mCTAR exercise and traditional dysphagia treatment (n = 12) | Control group, only traditional (n = 13) | Male: 6/6 | Type of stroke—haemorrhage: 5/7 | Side of stroke (right/left): 5/7/4/9 | Facial palsy: 1/1 | Dysarthria: 1/0 | | |
| **Diagnosis:** Stroke | Stroke diagnosis, Liquid aspiration | | | | | | | | |
| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
| **OD:** Stroke related dysphagia, hypopharyngeal residue found by VFSS | Intervention, modified jaw opening exercise (MJOE) (n = 6) | Control, isometric jaw closing exercise (n = 6) | Male: 5/5 | Post-onset weeks, mean (SD): 6.7 (2.1)/9.2 (4.0) | FOIS, n, Level 5/Level 6: 33/42 | | | | |
| **Diagnosis:** Stroke | | | | | | | | | |
| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
| **OD:** Stroke-related dysphagia and/or penetration on VFSS, nasogastric tube, or pen| Experimental/Control | Experimental/Control | Experimental/Control | Experimental/Control | Experimental/Control | Experimental/Control | Experimental/Control | Experimental/Control |
| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
| **OD:** Dysphagia determined by endoscopic swallowing evaluation | Experimental/Control | Experimental/Control | Experimental/Control | Experimental/Control | Experimental/Control | Experimental/Control | Experimental/Control | Experimental/Control |
| **Diagnosis:** Elderly patients with moderate-to-severe dysphagia. Diagnosis: NR | Study, original dysphagia treatment (n = 30) | Control (n = 30) | Male: 57/36 | | | | | |
| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
| **OD:** Dysphagia determined by endoscopic swallowing evaluation | Study/Control | Study/Control | Study/Control | Study/Control | Study/Control | Study/Control | Study/Control | Study/Control |
| **Diagnosis:** Elderly patients with moderate-to-severe dysphagia. Diagnosis: NR | Study, original dysphagia treatment (n = 30) | Control (n = 30) | Male: 57/36 | | | | | |
| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
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| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
| **OD:** Stroke related dysphagia, hypopharyngeal residue found by VFSS | Study/Control | Study/Control | Study/Control | Study/Control | Study/Control | Study/Control | Study/Control | Study/Control |
| **Diagnosis:** Stroke | | | | | | | | | |
| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
| **OD:** Stroke related dysphagia, hypopharyngeal residue found by VFSS | Study/Control | Study/Control | Study/Control | Study/Control | Study/Control | Study/Control | Study/Control | Study/Control |
| **Diagnosis:** Stroke | | | | | | | | | |
| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
| **OD:** Stroke related dysphagia, hypopharyngeal residue found by VFSS | | | | | | | | | |
| **Diagnosis:** Stroke | | | | | | | | | |
| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
| **OD:** Stroke related dysphagia, hypopharyngeal residue found by VFSS | | | | | | | | | |
| **Diagnosis:** Stroke | | | | | | | | | |
| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
| **OD:** Stroke related dysphagia, hypopharyngeal residue found by VFSS | | | | | | | | | |
| **Diagnosis:** Stroke | | | | | | | | | |
| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
| **OD:** Stroke related dysphagia, hypopharyngeal residue found by VFSS | | | | | | | | | |
| **Diagnosis:** Stroke | | | | | | | | | |
| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
| **OD:** Stroke related dysphagia, hypopharyngeal residue found by VFSS | | | | | | | | | |
| **Diagnosis:** Stroke | | | | | | | | | |
| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
| **OD:** Stroke related dysphagia, hypopharyngeal residue found by VFSS | | | | | | | | | |
| **Diagnosis:** Stroke | | | | | | | | | |
| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
| **OD:** Stroke related dysphagia, hypopharyngeal residue found by VFSS | | | | | | | | | |
| **Diagnosis:** Stroke | | | | | | | | | |
| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
| **OD:** Stroke related dysphagia, hypopharyngeal residue found by VFSS | | | | | | | | | |
| **Diagnosis:** Stroke | | | | | | | | | |
| **Inclusion:** | | | | | | | | | |
| **Exclusion:** | | | | | | | | | |
| Logemann, et al. [38] | USA |
|----------------------|-----|
| OD: Speech pathologist referral after swallow screening, patient aspirating thin liquids. |
| Diagnosis: Physician’s diagnosis of dementia or PD. Bedford Alzheimer Nursing Severity Scale; neurololgist rated PD using Hoehn and Yahr scale. |
| Inclusion: 50–95 yrs |
| Exclusion: Inability to perform chin down intervention |
| n = 742 |
| All patients received all 3 interventions (random order): |
| Chin-down intervention |
| Nectar |
| Honey-thickened liquids |
| Mild 0–3: 8 (13%) |
| Moderate 4–7: 35 (56%) |
| Severe 8–9: 19 (31%) |
| Age range: 50–79, 41% |
| Age range: 80–95, 59% |
| Male: 70% |
| PD-No dementia: 32% |
| PD-Dementia: 19% |
| Dementia–Other: 19% |
| Dementia–Single or multistroke: 15% |
| Dementia–Alzheimer’s: 15% |

| Manor, et al. [39] | UK |
|-------------------|----|
| OD: Referred to speech pathologist for evaluation of swallowing disturbances, confirmed via FEES. |
| Diagnosis: PD had been diagnosed according to the UK Brain Bank criteria |
| Inclusion: Diagnosis as above |
| Exclusion: History of other uncontrolled neurological or medical disorders interfering with swallowing |
| n = 42 |
| Experimental group - received video-assisted swallowing therapy (VAST, n = 21) |
| Control group - conventional therapy (n = 21) |
| Vast/Conventional therapy |
| Age yrs: 67.8 (6.3)/69.9 (9.7) |
| Disease duration (years): 7.4 (4.7)/8.8 (5.7) |
| Disease severity (H&Y-1–5): 2.2 (0.8)/2.2 (0.8) |
| Swallowing disturbances questionnaire: 14.7 (5.8)/14.3 (7.2) |
| Fiberoptic endoscopic evaluation of swallowing: 0.7 (0.4)/0.6 (0.4) |

| Mepani, et al. [40] | USA |
|-------------------|----|
| OD: Post deglutitive dysphagia, pharyngeal phase dysphagia, VFSS to confirm |
| Diagnosis: Stroke or chemoradiation for head and neck cancer |
| Inclusion: Pharyngeal phase dysphagia, incomplete UES opening, and postdeglutitive aspiration, hypopharyngeal residue, able to comply with protocol, dysphagia with aspiration of at least 3 month duration |
| Exclusion: History of parotid surgical procedures excluded |
| n = 11 |
| Traditional swallowing therapy (n = 6) [Denoted as ‘Mepani et al. (2009a)’ in Figure 4.] |
| Shaker Exercise (n = 5) [Denoted as ‘Mepani et al. (2009b)’ in Figure 4.] |
| Traditional/Shaker |
| Age years: 70.5 (9.5)/64 (22.8) |
| Male: 5 (83%)/3 (60%) |
| Etiology of dysphagia: |
| - CVA: 4 (67%) |
| - Cancer: 2 (33%) |
| - CVA: 4 (67%) |
| TPSS/Control |
| Age years: 62.0 (4.2)/63.5 (6.1) |
| Male: 3/4 |
| Stroke type (ischemic/hemorrhagic): 6/2/6/2 |
| Poststroke duration days: 56.0 (17.4)/59.9 (20.0) |
| MMSE: 22.87 ± 2.47 23.50 ± 2.00 |

| Moon, et al. [41] | Korea |
|------------------|------|
| OD: Aspiration or penetration, oropharyngeal residue, confirmed VFSS. |
| Diagnosis: Stroke or chemoradiation for head and neck cancer |
| Inclusion: Subacute stage 3–12 weeks after the onset of stroke |
| Exclusion: History of parotid surgical procedures excluded |
| n = 16 |
| TPSST plus traditional dysphagia therapy (n = 8) [Denoted as ‘Moon et al. (2018a)’ in Figure 4.] |
| Control, traditional dysphagia therapy (n = 8). [Denoted as ‘Moon et al. (2018b)’ in Figure 4.] |
| TPSST/Control |
| Age years: 64.3 (10.7)/65.8 (11.3) |
| Male n = 8/6 |
| Time since onset weeks: 27.4, 6 (6.3)/26.6 (6.8) |

| Park, et al. [42] | Korea |
|------------------|------|
| OD: Dysphagia confirmed by VFSS |
| Diagnosis: Stroke |
| Inclusion: Onset within 6 months; score ≥24 on the MMSE |
| Exclusion: Stroke prior to that resulting in dysphagia, severe orofacial pain, significant malocclusion or facial asymmetry, unstable breathing or pulse, tracheostomy, severe communication disorder, inadequate lip closure |
| n = 27 |
| Experimental group, Expiratory muscle strength training (EMST) (n = 14) |
| Placebo sham (n = 13) |
| Experimental/Placebo |
| Age years: 64.3 (10.7)/65.8 (11.3) |
| Male n = 8/6 |
| Time since onset weeks: 27.4, 6 (6.3)/26.6 (6.8) |

| Park, et al. [43] | Korea |
|------------------|------|
| OD: Dysphagia following stroke was confirmed by VFSS |
| Diagnosis: Stroke |
| Inclusion: Onset duration was <12 months, swallow voluntarily, MMSE score ≥20 |
| Exclusion: Secondary stroke, severe communication disorder, pain in the neck region, unstable medical conditions, head and neck cancer |
| n = 22 |
| Experimental, chin tuck against resistance exercise (CTAR; n = 11) [Denoted as ‘Park et al. (2018a)’ in Figure 4.] |
| Control group, only conventional dysphagia treatment (n = 11). [Denoted as ‘Park et al. (2018b)’ in Figure 4.] |
| Experimental/Control |
| Age years: 62.2 (17.3)/58.4 (12.5) |
| Male: 6/4 |
| Infarction: 7/6 |
| Time after stroke (weeks): 37.2 (54.3)/14 (14.4) |
| Oral feeding: 4/5 |
| Tube feeding: 7/6 |

| Park, et al. [44] | Korea |
|------------------|------|
| OD: OD after stroke by VFSS |
| Diagnosis: Stroke based on computed tomography or MRI |
| Inclusion: Inpatient, no significant cognitive problems (MMSE score ≥ 24) |
| n = 24 |
| Experimental, effortful swallowing training (EST; n = 12) [Denoted as ‘Park, Oh et al. (2019a)’ in Figure 4.] |
| Experimental/Control |
| Age years: 66.5 (9.5)/68.4 (11.2) |
| Male: 6/5 |
| Stroke lesion middle cerebral artery: 6/6 |
| OD: Dysphagia after stroke, by VFSS | Diagnosis: Stroke due to hemorrhage or infarction | n = 37 patients: |
|-----------------------------------|-----------------------------------------------|-----------------|
| Inclusion: <6 months of post-onset, nasogastric tube; absence of cognitive deficits. |
| Exclusion: Secondary stroke, difficulty in using both upper limbs, significant malocclusion or facial asymmetry, pain in the disc and cervical spine, limitations in opening jaw, use of cervical spine orthosis, trachoeotomy, severe communication difficulties associated with dementia or aphasia, presence of gastronomy tube, problems with the oesophageal phase of dysphagia |

| OD: Dysphagia screening-at least one severe symptom, validated in Greek Okhuma questionnaire |
| Diagnosis: Hemiaparesis following stroke |
| n = 70: |
| Inclusion: Hemiaparesis following stroke, at least one severe symptom of the validated Greek Okhuma questionnaire |
| Exclusion: Exclusion-Barthel Index >20, Motor Function Hemispheric Stroke Scale >25, history of OD. |

| OD: ‘Swallowing difficulties’ determined with Turkish version of the eating assessment tool (T-EAT-10) |
| Diagnosis: No neurological problems after neurologist’s examination |
| n = 50: |
| Inclusion: Over 65 yrs, adequate cognitive status. |
| Exclusion: Head/neck conditions affecting swallowing |

| OD: Dysphagia post stroke (VFSS) |
| Diagnosis: Recent stroke (4-20 wks) |
| n = 14: |
| Inclusion: Recent stroke, one repetition maximum posterior maximum isometric tongue-palate pressure measure ≤40 kPa at intake, stage transition duration if ≤350 ms on at least one liquid barium swallow at intake VFSS |
| Exclusion: Severe dysphagia with no functional opening of upper esophageal sphincter; pre-existing dysphagia or diagnoses of head and neck. |

| OD: Dysphagia based on DYMUS questionnaire (patient self-report) |
| Diagnosis: Nasopharyngeal carcinoma (NPC) patients after radiotherapy |
| n = 43: |
| Inclusion: Diagnosed as above |
| Exclusion: Dysphagia or trismus as initial symptoms of NPC excluded |

| OD: Pharyngeal dysphagia confirmed through VFSS |
| Diagnosis: Diagnosed as having stroke |
| n = 12. |
| Inclusion: Within 6 months post-onset, nasogastric tube; absence of cognitive deficiencies. |
| Exclusion: Secondary stroke, presence of other neurological, pain in the disc and cervical spine, cervical spine orthosis, presence of gastronomy tube, problems with the oesophageal phase of dysphagia |

| OD: Saliva swallowing (n = 12) [Denoted as ‘Park, Oh et al. (2019b)’ in Figure 4] |
| Time since stroke onset, wks: 24.4 (8.6)/25.7 (6.3) |

| OD: Saliva swallowing (n = 12) |
| Control, saliva swallowing (n = 12). [Denoted as ‘Park, Oh et al. (2019b)’ in Figure 4] |

| OD: Saliva swallowing (n = 12) |
| Time since stroke onset, wks: 24.4 (8.6)/25.7 (6.3) |

| OD: Saliva swallowing (n = 12) |
| Control, saliva swallowing (n = 12). [Denoted as ‘Park, Oh et al. (2019b)’ in Figure 4] |

| OD: Saliva swallowing (n = 12) |
| Time since stroke onset, wks: 24.4 (8.6)/25.7 (6.3) |

| OD: Saliva swallowing (n = 12) |
| Control, saliva swallowing (n = 12). [Denoted as ‘Park, Oh et al. (2019b)’ in Figure 4] |

| OD: Saliva swallowing (n = 12) |
| Time since stroke onset, wks: 24.4 (8.6)/25.7 (6.3) |

| OD: Saliva swallowing (n = 12) |
| Control, saliva swallowing (n = 12). [Denoted as ‘Park, Oh et al. (2019b)’ in Figure 4] |

| OD: Saliva swallowing (n = 12) |
| Time since stroke onset, wks: 24.4 (8.6)/25.7 (6.3) |

| OD: Saliva swallowing (n = 12) |
| Control, saliva swallowing (n = 12). [Denoted as ‘Park, Oh et al. (2019b)’ in Figure 4] |

| OD: Saliva swallowing (n = 12) |
| Time since stroke onset, wks: 24.4 (8.6)/25.7 (6.3) |

| OD: Saliva swallowing (n = 12) |
| Control, saliva swallowing (n = 12). [Denoted as ‘Park, Oh et al. (2019b)’ in Figure 4] |

| OD: Saliva swallowing (n = 12) |
| Time since stroke onset, wks: 24.4 (8.6)/25.7 (6.3) |
| Country | Diagnosis | Inclusion | Exclusion | Intervention/Control | Age years | Male: | Disease Duration (years) | Expanded Disability Status Scale | MS Type-Relapse-Remitting: | MS Type-Primary Progressive | MS Type-Secondary Progressive |
|---------|-----------|-----------|-----------|---------------------|-----------|-------|-------------------------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|
| Iran    | Established diagnosis of MS according to McDonald’s criteria | - | - | Experimental (TDT), sensorimotor exercises and swallowing manoeuvres (n = 10) [Denoted as ‘Tarameshlu et al. (2019a)’ in Figure 4.] | 47.5 (12.9)/39.9 (9.7) | 25/22 | 6.8 (2.9)/6.1 (2.7) | 3.6(2.1)/3.2(2.5) | 4/7 | 4/1 | 2/2 |

| USA     | Diagnosis: PD-diagnostic criteria of the UK Brain Bank | Inclusion: 55–85 yrs, same PD medication, >24 MMSE. Exclusion: other neurologic disorders; head/neck cancer | - | Expiratory muscle strength training (EMST; n = 30) | 66.7 (8.9)/68.5 (10.3) | 25/22 | Male: 66.7 (8.9)/68.5 (10.3) | 81 (9)/81 (21) | 24 |
|---------|-----------|-----------|-----------|---------------------|-----------|-------|-------------------------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|

| Japan   | Diagnosis: NR (Community-dwelling, ≥65 yrs) | Inclusion: Receiving long-term care via day-service or day-care program, mild cognitive impairment/dementia | Exclusion: Severe or moderate dementia, inability to perform training | Intervention/Control | 80 (7)/79 (7) | Male: 19/28 | Tongue pressure (kPa): 23.3 (8.3)/23.3 (10.0) | EAT-10, median (IQR): 7 (5–13)/8 (4–11) | 35 |

| France  | Diagnosis: NR. (Sitting abnormality- by seated postural control measure, SPCM). | Inclusion: >18 years; DHI score >1, score >0 on 1 item SPCM, chronic dysphagia. | Exclusion: NR | D-/D+ | Age years (total sample): 61.5 (11.8) | Male, n (total sample): 25 | Degenerative dysphagia, N (total sample): 24 | NIHSS: 1.3 (1.4)/1.3 (1.6) | 1.7 (1.3)/1.9 (1.9) | 6.0 (0.9)/5.8 (1.1) | 2/2 |

| China   | Diagnosis: Stroke | Inclusion: Swallowing dysfunction | exclusion: NR (admitted patients with dysphagia) | Intervention, nursing intervention (n = 60) | Age years: 70.6 (7.4)/70.3 (7.4) | Male: 33/32 | Control, conventional nursing service (n = 60) | 60:

Notes: ABI = Acquired brain injury; BDI = Beck Depression Inventory; BDST = Burke Dysphagia Screening Test; CVA = cerebrovascular accident; DOSS = Dysphagia Outcome and Severity scale; FEES = Fiberoptic Endoscopic Evaluation of Swallowing; FOIS = Functional Oral Intake Scale; GCSC = Glasgow Coma Scale; H&Y disability score = Hoehn and Yahr disability score; K-MMSE or MMSE-K = Mini-mental examination Korean version; K-NIHSS = Korean version of National Institute of Health Stroke Scale; MASA = Mann Assessment of Swallowing Ability; MBS = Modified Barium Swallow; MIE = Minimally Invasive Oesophagectomy; MDTTP = McNeill Dysphagia Therapy Program; MMSE = Mini-Mental State Examination; MOCA = Montreal Cognitive Assessment; NIHSS = National Institute of Health Stroke Scale; NMES = Neuromuscular Electrical stimulation; NR = Not reported; OD = Oropharyngeal dysphagia; PAS = Penetration-Aspiration Scale; PD = Parkinson’s disease; P-DHI = Persian Dysphagia Handicap Index; PDQ-39: Parkinson’s Disease Questionnaire-39; PNF = proprioceptive neuromuscular facilitation; RCT = Randomised Controlled Trial; SLP: Speech-Language Pathology; SRS = Swallowing Rating Scale; SSA = Standardized Swallowing Assessment; SIS-6 = Swallowing Impairment Score; SWAL-QOL = Swallow Quality-of-Life Questionnaire; TDCS = transcranial Direct Current Stimulation; UC = Usual Care; VDS = Video-fluoroscopic Dysphagia Scale; VFSS = Video-Fluoroscopic Swallowing Study; WHO = World Health Organisation; WST = Water Swallow Test; TWST = Timed Water-Swallow Test.
Experimental group showed significant improvement in clinical evaluation of dysphagia compared to two other groups regarding solid ($p < 0.001$) and liquid ($p = 0.022$). Analysis of FEES did not show differences between groups. Experimental group presented with significant improvement in scores of domains frequency of symptoms ($p = 0.029$) and mental health ($p = 0.004$) on the SWAL-QOL when compared with the groups that did not receive intervention.

### Table 3. Outcome of behavioural interventions for people with oropharyngeal dysphagia.

| Study | Intervention Goal | Intervention Agent, Delivery and Dosage | Materials and Procedures | Outcome Measures | Treatment Outcome |
|-------|-------------------|----------------------------------------|--------------------------|-----------------|------------------|
| Ayres et al. [19] | To verify the effectiveness of a manoeuvre application in swallowing therapy in patients with PD. | Three groups: Experimental group: Chin-down posture manoeuvre (patient instructed to ‘swallow lowering the head until chin touches in the neck’). Patients performed manoeuvre twice a day, swallowing saliva, during meals, throughout the week, at home. Patients were given a form to record the number of times the manoeuvre was performed at home. Patients also given instructions for optimal feeding and swallowing related to ‘swallowing orientations’: (1) environment during feeding (2) posture (3) meal-time (4) oral hygiene. Written instructions given. Control group: No intervention received during 4-week period. Written instructions given. | Primary outcomes: FEES; Clinical evaluation (checking 21 signs and symptoms of oropharyngeal dysphagia and rating these as present or absent); FOIS; SWAL-QOL. | Experimental group showed significant improvement in clinical evaluation of dysphagia compared to two other groups regarding solid ($p < 0.001$) and liquid ($p = 0.022$). Analysis of FEES did not show differences between groups. Experimental group presented with significant improvement in scores of domains frequency of symptoms ($p = 0.029$) and mental health ($p = 0.004$) on the SWAL-QOL when compared with the groups that did not receive intervention. |
| Carnaby et al. [19] | Compare standard low-intensity and high-intensity behaviour interventions with usual care (UC) for dysphagia | | | |
| Carnaby et al. [20] | Effectiveness and safety of exercise based swallowing therapy and neuromuscular electrical stimulation for dysphagia | | | |
| Carnaby et al. [21] | | | | |
| Study | Intervention/Exercise | Dosage | Primary Outcomes | Experimental Group | Control Group |
|-------|-----------------------|--------|------------------|-------------------|---------------|
| Choi et al. [22] | Effects of Shaker exercise on aspiration and oral diet | **Intervention agent:** Caregiver (SE), occupational therapist (CDT)  
**Dosage:** 30 min/day, 5 days/wk × 4 wks | **Shaker Exercise (SE):** Isometric and isokinetic movements. 3 head lifts held for 60 s in supine; 60 s rest. 30 reps head lifts observe toes without raising shoulders-without hold.  
**Conventional Dysphagia Therapy (CDT):** Orofacial muscle exercises, thermal tactile stimulation, therapeutic/compensatory manoeuvres. | Experimental group greater lifts held for 60 s in supine; 60 s rest. 30 reps head lifts observe improvement on PAS (*p* < 0.05)  
No significance between control groups for time until end of meal. |  |
| DePippo et al. [23] | Effect of graded intervention on occurrence of dysphagia related complications | **Intervention agent:** Dysphagia therapist (SLP)  
**Dosage:** Bi-weekly session monitoring for all groups | Group A—Patient-managed diet. One session-therapist recommended diet based on MBS results and compensatory swallowing techniques. Patient chose diet (regular vs. graded).  
Group B—Therapist-prescribed diet (MBS) and swallowing techniques, evaluated every other week.  
Group C—Therapist prescribed diet and daily reinforcement of swallowing techniques through mealtime dysphagia group. | Primary outcomes: Dysphagia related complications: Pneumonia, dehydration, calorie-nitrogen deficit, recurrent upper airway obstruction, and death. |  |
| Eom et al. [24] | Effect of resistance Expiratory Muscle Strength Training (EMST) on swallowing function | **Intervention agent:** NR  
**Dosage:** 5 days p/wk × 4 wks, 5 sets of 5 breaths on device × 25 p/day. Both groups treatment 30 min × 5 days/wk × 4 wk | Experimental group (EMST + Conventional treatment): Portal Emphoratory Muscle Strength Trainer (EMST150). Patients opened mouth after inhalation, EMST mouthpiece between lips. Blew strongly and rapidly until pressure release valve within EMST device opens. Pressure release set to open if pressure target exceeded. < 1-min break after each session, for muscle fatigue and dizziness.  
Placebo group (Sham EMST + Conventional treatment): Trained using a sham non-functional EMST device with no loading device. Conventional treatment. | Primary outcomes: VDS and PAS based on a VFSS to analyse oropharyngeal swallowing function. | Experimental significant in VDS pharyngeal phase (*p* = 0.02 and 0.01) and PAS vs. placebo (*p* = 0.01). Both significant VDS and PAS in all phases (*p < 0.05*). Experimental only significant in PAS (*p = 0.01* vs. 0.102). |
| Gao and Zhang [25] | Effects of rehabilitation training on dysphagia and psychological state | **Intervention agent:** NR  
**Dosage:** 3 sessions/actions performed morning, midday and evening. 7 days p/wk × 42 days | All patients received routine treatment including internal medicine, traditional rehabilitation and routine nursing.  
Control: Traditional tongue and mouth exercises. Each movement repeated 10 times as one session.  
Shaker exercise: Supine position, single action raised head to look at feet. 30 reps = set of actions. Perform 3 sets of actions—continuously or with 1-min relaxation until complete. (Denoted as ‘Gao & Zhang, 2017a’ in Figure 5.)  
Chin Tuck Against Resistance (CTAR) exercise: Patients seated tucking chin to compress inflatable rubber ball for 30 reps = set of actions. Perform 3 sets, continuously or with relaxation. (Denoted as ‘Gao & Zhang, 2017b’ in Figure 5.) | Primary outcomes: Dysphagia: VFSS at baseline, 2, 4, 6 wks post. Swallowing function, PAS Psychological state: Self-Rating Depression Scale (SDS) baseline, 6 wks post. | Degrees of dysphagia improvement, between 2–4 wks in CTAR and Shaker. Significantly higher in CTAR (87%) and Shaker (77%) vs. control (43%) (all *p < 0.05*). Significantly lower SDS in CTAR vs. Shaker/control 6 wks post (all *p < 0.05*). |
| Guillén-Solà et al. [26] | Effectiveness of inspiratory/expiratory muscle training (IEMT) and neuromuscular electrical stimulation (NMES) | **Intervention agent:** Occupational, speech, physical therapist  
**Dosage:** Control: 3 hrs p/day × 5 days wk × 3 wks.  
**Control/SST:** Multidisciplinary inpatient rehabilitation for mobility, activities of daily living, swallowing and communication. Education self-management of dysphagia, oral exercises and compensatory techniques based on VFSS.  
**EMST + SST:** Inspiratory/Expiratory Muscle Training (EMST)-respiratory training, 5 sets of 10 respirations, 1 min unloaded | Primary outcomes: Dysphagia severity by PAS. Respiratory muscle strength (maximal inspiratory and expiratory pressures). Post- and 3-month follow-up. | Maximal respiratory pressures most improved Group 2: treatment effect 12.9 (CI 4.5–21.2) and 19.3 (CI 8.5–30.3) for maximal inspiratory and expiratory pressures. Swallowing security improved. |  |
| Study | Intervention | Outcome Measures | Results |
|-------|--------------|-----------------|---------|
| Haggled et al. [27] | Group A: Orofacial sensory-vibration stimulation; Group B: Orofacial sensory-vibration stimulation + oral neuromuscular training | Swallowing rate (timed water swallow test) | Significant improvement in Group B compared to Group A (p < 0.001) |
| | | Lip force | Significant improvement in Group B compared to Group A (p < 0.0001) in TWST, but no significant between-group difference in swallowing rate. |
| | | Changes in swallowing rate as measured by the Timed Water Swallow Test (TWST) | Changes in lip force as measured by lip-force test + swallowing dysfunction measured by VFS (in lateral projection). |
| Hwang et al. [29] | Effect of tongue stretching exercises (TSE) on tongue motility and oromotor function in patients with dysphagia after stroke. | Primary outcomes: Oromotor function—Oral phase events of VDS, VFS | Lip force: Significant improvement in lip force in Group 2 (p < 0.001) compared to non-significant improvement in Group 1 (p = 0.079). Improvement in Group 2 maintained at 12 month follow up. |
Intervention feasible. PAS and FOIS improved in both groups, no group differences. Swallowing specific parameters reflected clinically observed changes.

### Primary outcomes:
- FOIS, PAS, and electrophysiological swallowing specific parameters

### Experimental treatment:
- Facial Oral Tract Therapy (F.O.T.T.) concept-rehabilitation intervention using structured tactile input and nonverbal facilitation techniques (to allow for effective function in meaningful daily activities).

#### Control group:
- Treatment comprised stimulating activities in the conventional swallowing rehabilitation. Inspiration-Expiration pressure 15–20 cm H2O, increased to 40 cm H2O for 2 s.
- Expiration—similar pressure 10–20 cm H2O above positive pressure; held 3–6 s, simulating airflow during cough. Patient coordinated respiratory rhythm to cough assist machine.

#### Study group
- **MIE exercise:**
  - MIE-100 Cough Assist
  - Conventional dysphagia rehabilitation of oral motor and sensory stimulation, NMES, oral exercises for safe swallow.

#### Intervention agent:
- **NR**

### Dosage:
- 20 sessions once a day, 3 × wk × 2 wks.

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### Jang et al. [31]

Effects of Mechanical Inspiration and Expiration (MIE) exercise using mechanical cough assist on velopharyngeal incompetence

#### Intervention agent:
- NR

#### Dosage:
- 20 sessions
- Both groups, 30 min 2 × day; 5 × wk × 2 wks.

### To investigate the effects of NMES plus upper spine cervical mobilisation on forward head posture, and swallowing in stroke patients with dysphagia.

#### Intervention agent:
- Joint mobilisation was performed by a physical therapist (with over 160 h of manual therapy education).

#### NMES was delivered by 3 experienced OTs.

#### Dosage:
- once a day, 3 × times a week, for 4 weeks;
- both groups received NMES for 30 min;
- experimental group received 10 min of upper cervical spine mobilisation; control group received 10 min of sham mobilisation.

### All interventions were performed in sitting position.

#### NMES:
- **Intervention group** received upper cervical spine (C1–2) mobilisation with NMES.
- **Mobilisation:** Therapist used one hand to hold the subject’s C1 (atlas); other hand placed on subject’s occiput. Mobilisation force could not be standardised. NMES was applied to the suprahyoid using VitalStim®. Electrodes attached to the motor point of the suprahyoid muscles (digastric) to induce anterior excursion and vertical elevation movements of hyoid bone during normal swallowing. Stimulation was applied by gradually increasing the intensity to the level that patients felt a grabbing sensation in the neck without pain or laryngospasm.

#### Control group:
- Patients received upper cervical spine sham mobilisation combined with NMES.

### Primary outcome:
- Forward head posture measured by CCFT (Stabilizer™ Pressure Biofeedback) and craniovertebral angle (CVA).

#### Swallowing function measured by VFS and PAS.

### The intervention group showed significantly better scores in CCFT (p < 0.05) and in CVA (p < 0.05) than in control group. PAS scores were significantly better in the intervention group compared to control group (p < 0.05).

### Significant increase in VFS total score and PAS than in the control group (p < 0.05).

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### Jeon et al. [32]

The effects of Proprioceptive Neuromuscular Facilitation (PNF) on swallowing function of stroke pts with dysphagia

#### Intervention agent:
- NR

#### Dosage:
- PNF-based short neck exercises 3 times a week for 30 min each time for 6 weeks

### Experimental group: PNF

1. Patients started by lying on a bed with head and neck positioned off the bed (tester supported left laryngeal region with his right hand and placed left fingertips below patient’s jaw)
2. Patient instructed to look at target object in a direction 15 degrees diagonally to the right side
3. Tester then initiated given exercises by moving the patient’s neck in a diagonal direction opposite to the direction specified

### Primary outcome:
- New VFS and ASHA NOMS Scales.

### Statistically significant improvements in: premature bolus loss, residue in the valleculae, laryngeal evaluation, epiglottic closure, residue in pyriform sinuses, coating of pharyngeal wall after swallowing, pharyngeal transit time and aspiration on both new VFS scale and
| Study | Intervention | Description | Dosage | Primary outcomes | Secondary outcomes |
|-------|--------------|-------------|--------|------------------|--------------------|
| Kim and Park [34] | Intervention agent: Occupational therapist | mCTAR exercise | 30 min × 5 days a week, for 6 weeks | Primary outcomes: Aspiration and oral diet -PAS and FOIS. | No temporomandibular joint or neck pain. Intervention group, DMH decrease where anterior HD ended and an increase in anterior HD were seen. Control, no changes. |
| Koyama et al. [35] | Intervention agent: Speech pathologist/physician | MJOE | 4 × sets daily, 5 × p/wk × 6 wks. (6 s × 5 reps = 1 set) | Primary outcomes: VFSS was performed before and after exercise. The distance between the mental spine and the hyoid bone (DMH) and hyoid displacement (HD) were measured. | No temporomandibular joint or neck pain. Intervention group, DMH decrease where anterior HD ended and an increase in anterior HD were seen. Control, no changes. |
| Krajczy et al. [36] | Intervention agent: Physiotherapist | Physiotherapy program average 60 min × day, × 15 days | Control/both groups: Safe food education and neurological physiotherapy depending on patient dysfunction. Therapy included passive, assisted, supported and respiration exercises, erect posture, walking re-education, and training on NDT Bobath and PNF methods. Study group: + original dysphagia treatment, restoring chewing and swallowing functionality—Strengthening and breathing exercises and thermal stimulation. | Primary outcomes: Swallowing function - Timed test of swallowing – Controlled swallowing after swallowing blended food. Reflex categorised as good or delayed. | All Statistically significant differences between groups after therapy (p < 0.01). |
| Kyodo et al. [37] | Intervention agent: Gastroenterologists | To evaluate the effectiveness of puree diets containing a gelling agent for the prevention of aspiration pneumonia in elderly | Patients underwent endoscopic swallowing evaluation while sitting in a chair/sitting up in bed. Images of oropharynx and larynx were displayed on a monitor and recorded on digital video recorder. Pureed diet without gelling agent was made by mixing 100 g of white rice and 50 mL of water with a blender for one minute. | Primary outcome: Presence of material in throat using endoscopic cyclic ingestion score (0 to 4) | Residuals in throat were significantly less likely with pureed rice with than without the gelling agent (median cyclic ingestion score (range): 1 (0–4) vs. 2 (0–4); p = 0.001. |
Irrespective of presence or absence of the gelling agent, the sense of materials in the throat was significantly less frequent in older patients \( (p < 0.01) \). No adverse events occurred.

| Intervention | Dosage (average): | Primary outcomes: |
|--------------|-------------------|-------------------|
| Logemann et al. \[38\] | 3 treatments for aspiration on thin liquids—chin-down posture, nectar-thickened liquids, or honey-thickened Liquids. | Swallowing function-VFSS |
| Control–conventional therapy: Both interventions swallowing exercises and compensatory therapy based on FEES. Compensatory strategies carried out with different food and liquid consistencies in clinic, patient practiced at home. VAST: video-assisted tool during each session, for educating and assisting understanding structure of swallowing. Patients observed a normal swallowing process and their distorted one. After learning compensatory technique, patient practiced it during drinking and eating in the clinic after observing video then at home. During next four sessions patients observed video feeding by questioners with suitable compensatory swallowing technique while eating and drinking focusing on the new swallowing behaviour. | Significant improvement in swallowing functions both groups. FEES significantly greater reduction in food residues in pharynx in VAST vs. conventional treatment group. SWAL-QOL scores significant between groups favour of VAST: burden, eating desire, social functioning, mental health, symptom frequency \( (p < 0.01) \). |
| Manor et al. \[39\] | Effectiveness of visual information while treating swallowing disturbances in patients with PD. | Swallowing function-by fiberoptic endoscopic evaluation of swallowing (FEES). Quality of life—quality of life and degree of pleasure from eating assessed by questioners |
| Traditional therapy: 5 times daily. Laryngeal and tongue ROM exercises and swallowing manoeuvres (Super-Supraglottic Swallow, Mendelsohn Manoeuvre, Effortful Swallow). Shaker Exercise: 3 times per day for 6 weeks. Isometric and isotonic head-lift in supine position. Patients raised head high and forward to observe toes. Isometric–3 times head lifts held 60 Videofluoroscopy s, 60-s rest period. Isokinetic–30 head lifts at constant velocity, performed without holding or rest periods. | After therapy, the percent change in thyrohyoid distance in the Shaker Exercise group was significantly greater vs. traditional therapy \( p = 0.034 \). |
| Mepani et al. \[40\] | Effect of the Shaker exercise on thyrohyoid muscle Shortening improve pharyngeal dysphagia | Change in thyrohyoid muscle shortening by |
| Traditional dysphagia therapy: thermal tactile stimulation, Mendelsohn manoeuvre, effortful swallow, diet modification. | |
| Moon et al. \[41\] | Effects of Tongue pressure strength and accuracy training (TPSAT) on tongue pressure strength, swallowing function, and swallowing. Discipline who created gelling agent (intervention) NR. | Tongue pressure strength - maximum isometric tongue pressure (MIPs) of anterior, TPSAT with traditional dysphagia significantly improved MASA, SWAL-QOL, and MIPs. Traditional dysphagia significantly |
quality of life in stroke patients with dysphagia. Traditional therapy performed 30 min × twice daily. Both groups, daily 5× times wk × 8 wks.

TPSAT with traditional dysphagia treatment: TPSAT consisted of an anterior and posterior isometric tongue strength exercise and an isometric tongue accuracy exercise. The protocol involved five sets of tongue-to-palate presses, 6 reps per set for each session. Isometric tongue accuracy exercise, amplitudes were set at 50, 75, 100% of maximum pressure from first isometric strength. Participants generated precise pressures within 10 kPa error for each amplitude.

Park et al. [42]

Effects of EMST on the activity of suprahyoid muscles, aspiration and dietary stages in stroke patients with dysphagia.

Intervention agent: Occupational therapist
Dosage: 5 days × wk × 4 wks. 5 sets × 5 breaths on device, 25 breaths per day.

Experimental group: resistance set at 70% range of MEP (Maximal Primary outcomes: Expiratory Pressure). Subjects open mouth following maximum inhalation, EMST mouthpiece between lips, close mouth. Blow strong and fast until pressure release valve in EMST device opens- expiratory pressure exceeded set target. Placebo group: training using sham device-non-functional device, little effect of physiologic load on targeted muscles.

Experimental significantly more in suprahypoid muscle activity (p < 0.01), liquid PAS (p = 0.03) and FOIS (p < 0.06), but not semisolid type PAS (p = 0.32), vs. placebo.

Park et al. [43]

Effect of chin tuck against resistance exercise (CTAR) on the swallowing function in patients with dysphagia following subacute stroke.

Intervention agent: Occupational therapist
Dosage: 30 min × day, × 5/wk, × 4 wks

CTAR: Isometric CTAR, patients chin tuck against device 3 × 60 s no repetition. Isotonic CTAR, patient 30 reps by strongly pressing against resistance of the device and releasing it. Therapist demonstrated exercise methods.

Conventional dysphagia treatment: Both groups - orofacial muscle exercises, thermal tactile stimulation, and therapeutic or compensatory manoeuvres.

Primary outcomes: Swallowing function - Functional Dysphagia Scale (FDS) and PAS, based on VFSS

Experimental group: significant improvement in oral cavity, laryngeal elevation/epiglottic closure, residue in valleculae, and residue in pyriform sinuses of FDS and PAS compared vs. controls (p < 0.05, all).

Park et al. [44]

Effects of Effortful Swallowing Training (EST) on tongue strength and swallowing function in patients with stroke.

Intervention agent: Occupational therapist
Dosage: Training 30 min, 5× days per wk × 4 wks. Both groups conventional dysphagia treatment 30 min/day, 5 days/wk × 4 wks.

Experimental EST: Patients pushed tongue onto palate, squeezing neck muscles, swallow forcefully. Performed 10 times p/session, 3 sessions p/day. Effortful swallowing confirmed by therapist through visual observation and palpation.

Control group: Swallow naturally without intentional force. Patients given small spray of water to induce swallowing, and rest.

Both groups received conventional dysphagia therapy (compensatory techniques – chin tuck, head tilting, rotation; therapeutic techniques – orofacial muscle exercises, thermal tactile stimulation using ice sticks, expiratory training).

Primary outcomes: Tongue strength - Iowa Oral Performance Instrument. Oropharyngeal swallowing function VDS, based on VFSS

Experimental group greater improvements in anterior and posterior tongue strength vs. control (p = 0.05 and 0.04), and greater improvement in oral phases of VDS (p = 0.02).

Park et al. [45]

Effects of game-based Chin Tuck against resistance exercise (gbCTAR) and head-lift exercise on swallowing function and compliance in dysphagia post-stroke

Intervention agent: Occupational therapist
Dosage: 5 × wk × 4 weeks. Traditional dysphagia treatment (TDT) 30 min per day

Experimental group: performed gbCTAR exercise LES 100 device. Before gbCTAR exercise, 1-RM measured for resistance values. 1-RM, resistance bar placed directly beneath jaw, and chin tuck directed against resistance. gbCTAR exercise at threshold of 70% 1-RM, divided into isometric and isotonic exercises, combined with the game.

Control group: head lifts exercises in supine (isometric and isotonic).

Primary outcomes: Swallowing function-VDS and PAS.
Dietary assessment-FOIS Compliance with the 2 exercises- motivation,

No significant between group difference in VDS, PAS, FOIS. Compliance, motivation and interest Scores significantly higher, and scores for physical effort needed and fatigue
| Study | Intervention | Dosage | Primary Outcomes | Experimental group | Placebo group | Experimental vs. Placebo | Experimental vs. Control | **References** |
|-------|--------------|--------|------------------|-------------------|---------------|------------------------|-------------------------|----------------------|
| Park et al. [46] | Effect of Resistive Jaw Opening Exercise (RJOE) on hyoid bone movement, aspiration, and oral intake level in stroke patients. | **Intervention agent:** Occupational therapist  
**Dosage:** 30 min × 5 times wk × 4 wks. |  | Experimental group: RJOE device to provide resistance to suprahyoid muscles. Isometric exercise, 30 s with device resistors pressed downward (3 times, 30–60 s of rest). Isotonic exercise repeatedly depressed by RJOE by holding device resistance down for 2–3 s then returned to original state (10 reps, 3 sets) with 30 s rest.  
Placebo group: RJOE using 1-mm thick device with almost no resistance to suprahyoid muscles. Exercise type and frequency of RJOE same as experimental group. | Both groups received conventional dysphagia therapy after intervention, which involved orofacial muscle exercises, thermal tactile stimulation and therapeutic or compensatory manoeuvres. |  |  |  |  |
| Ploumis et al. [47] | Evaluate cervical isometric exercises in dysphagic patients with cervical spine alignment disorders due to hemiparesis after stroke. | **Intervention agent:** Allied health  
**Dosage:** inpatient 12 wks, speech 30 min daily. Experimental-4× reps 10 min, 3× day, 12 wks. |  | All patients-inpatient program including physiotherapy, occupational and speech therapy. Speech included deglutition muscle strengthening, compensatory techniques.  
Experimental group: plus cervical isometric strengthening exercises contract neck muscles under resistance forward-backward-sideways.  
Control group: Regular speech therapy plus sitting balance. |  |  |  |  |  |
| Sayaca et al. [48] | Whether combined isotonic technique of Proprioceptive Neuromuscular Facilitation (PNF) is superior to Shaker exercises in improving function of swallowing muscles. | **Intervention agent:** Shaker ‘CS’ (?). PNF physiotherapist  
**Dosage:** Each exercise set 1 x per day, 3x wk x 6 wks. |  | Shaker exercises: isometric (3 reps) and isotonic contractions (30 reps) neck flexor muscles. Patients raised head to observe toes without raising shoulders. Isometric- lifted head, held for 1 min 3 times, 1-min rest. Isotonic- lifted head 30 reps, no holding.  
PNF: Combined isotonic technique- concentric, stabilizing and eccentric contraction without relaxation. Stabilizing contractions to improve control, force, coordination, and eccentric contraction. Moved head against resistance with open mouth—kept position for 6 s against resistance in seated position; kept position while physiotherapist moved back to initial position. 30 reps per day. | T-EAT-10 decreased both groups (p < 0.001). Water swallowing capacity and volume improved both groups (p < 0.001). No change in swallowing speed both groups (p > 0.05). Maximal voluntary contraction of suprahyoid muscles higher in PNF vs. Shaker (p < 0.05). |  |  |  |  |
| Steele et al. [49] | Compare outcomes of two tongue resistance training protocols | **Intervention agent:** Speech pathologist  
**Dosage:** 24 sessions (TPPT or TPSAT), 2–3× wk, 8–12 wks. 60 tongue-pressure tasks per session. |  | Tongue-pressure profile training (TPPT): emphasized pressure-timing patterns that are typically seen in healthy swallows by focusing on gradual pressure release and saliva swallowing tasks.  
Tongue- pressure strength and accuracy training (TPSAT): emphasized strength and accuracy in tongue-palate pressure generation and did not include swallowing tasks. | Both groups significant tongue strength and post-swallow vallecular residue with thin liquids. Stage transition duration (bolus control), PAS no significant differences. |  |  |  |  |
| Study | Intervention | Dosage | Primary Outcomes | Secondary Outcomes |
|-------|-------------|--------|------------------|--------------------|
| Tang et al. [50] | Both groups routine treatment. Rehabilitation group: training by therapists at hospital, continued at home post-discharge by exercise booklet, guardian oversight and calendar Exercises: | 3× per day, each 15 cycles, 45 cycles per day. | Percentage of patients with effective results in rehabilitation higher than control (p = 0.02). Control IID significantly decreased at Post (p = 0.001), both groups increased at 3 months, rehabilitation group less than controls (p = 0.004). Trismus in rehabilitation higher vs. control (p = 0.02). | Trismus-LENT/SOMA score and the intercisor distance (IID). |
| Tarameshlu et al. [51] | Both groups routine treatment (e.g., anti inflammatory treatment for aspiration pneumonia) | 6 weeks, 18 sessions, 3 x per week, every other day. | Significant improvement in swallowing function. | Trismus-LENT/SOMA score and the intercisor distance (IID). |
| Troche et al. [52] | Both groups routine treatment (e.g., anti inflammatory treatment for aspiration pneumonia) | 4 weeks, 5 days per week, for 20 min per day, using a calibrated or sham, handheld device. | Significant improvement in swallowing function. | Trismus-LENT/SOMA score and the intercisor distance (IID). |
| Wakabayashi et al. [53] | Both groups routine treatment (e.g., anti inflammatory treatment for aspiration pneumonia) | 10 s; 1 set = 10 reps. 2 sets per day; 3× per wk × 3 months | Significant improvement in swallowing function. | Trismus-LENT/SOMA score and the intercisor distance (IID). |
| Woisard et al. [54] | Both groups routine treatment (e.g., anti inflammatory treatment for aspiration pneumonia) | 1 x training session with device (D+ group) | Significantly better posture both groups (p < 0.001), more hyoid bone motion in D+ group. Significant mean difference for D+ group vs. D− group | Trismus-LENT/SOMA score and the intercisor distance (IID). |
group) and without device (D- group).

**D+ group:** in charge to determine characteristics of the device required so they could have them during the training session. Instruction for patients was to put the personalised instructions into practice by using the device.

Measurement of hyoid bone movement during swallowing. VFSE and questionnaire.

**D- group:** for horizontal and vertical movement. Other swallowing markers not significant.

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**Zhang and Ju [55]**

Clinical improvement of nursing intervention in swallowing dysfunction of elderly stroke patients.

*Intervention agent:* Nursing staff

*Dosage:* NR

**Control group:** conventional nursing service that strictly conforms to the doctor’s advice.

**Nursing intervention:** (1) Psychological intervention, nurses communicate with patients/family, evaluates psychological state, encourages and comforts. (2) Health education, nurse introduces knowledge about swallowing dysfunction and effects through videos and images. (3) Rehabilitation exercises, pronunciation training, muscle training, mouth opening exercises, ingestion training. (4) Diet intervention, appropriate foods should be chosen according to specific conditions.

Primary outcomes: Swallowing dysfunction–30 mL water drink test

Living quality-assessment questionnaire of living quality (GQOL-74), includes physical, psychological and social functions, and material life.

Pulmonary infection–rate

Nursing satisfaction–self-made questionnaire.

*Improved swallowing dysfunction higher in intervention vs. control (p < 0.05). Scores of physical, psychological and social functions, and material life and nursing satisfaction higher in intervention vs. control (p < 0.05). Pulmonary infection lower in intervention vs. control (p < 0.05).*

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*Terminology as by authors. Notes. CVA = Cerebrovascular accident; EMST = Expiratory Muscle Strength Training; FEES = Fiberoptic Endoscopic Evaluation of Swallowing; FOIS = Functional Oral Intake Scale; MASA = Mann Assessment of Swallowing Ability; MBS = Modified Barium Swallow; MIE = Minimally Invasive Oesophagectomy; MDTP = McNeill Dysphagia Therapy Program; MEP = Maximum Expiratory Pressure; NMES = Neuromuscular Electrical stimulation; NR = Not reported; OD = Oropharyngeal dysphagia; PAS = Penetration-Aspiration Scale; PD = Parkinson’s disease; P-DHI = Persian Dysphagia Handicap Index; PNF = Proprioceptive Neuromuscular Facilitation; PRRS = Pharyngeal Residue Rating Scale; QoL = Quality of life; RCT = Randomised Controlled Trial; SIS-6 = Swallowing Impairment Score; SWAL-QOL = Swallow Quality-of-Life Questionnaire; VDS = Video-fluoroscopic dysphagia scale; VFSS = Video-Fluoroscopic Swallowing Study; TWST = Timed Water-Swallow Test; VDS = Videofluoroscopic Dysphagia Scale; VFSE = Videofluoroscopic Examination.*
Participants (Table 2). The 37 studies included a total of 2656 participants (mean = 72; SD = 124.5), with the sample sizes across studies ranging from 10 [30] to 742 participants [38]. All but two studies reported the mean age of participants [38,49], which was 65.6 years (SD = 8.8). Participant age range was only reported in five studies, ranging between 55 [36] and 95 [38] years. The mean percentage of male participants across all studies was 55.8% (SD = 13.7).

Most studies included stroke patients (n = 24). Other diagnoses included: patients with Parkinson’s disease [19,39,52], acquired brain injury [30], multiple sclerosis [51] and nasopharyngeal cancer [50]. Two studies included a mixed patient population with Parkinson’s disease or dementia [38], and stroke or head and neck cancer patients after chemoradiation [40]. Five studies did not provide further details on diagnoses [28,38,49,54,55]. The most frequent method for confirming OD was VFSS (n = 17), with only four studies using FEES (n = 4) [20,31,38,40]. Seven studies used non-instrumental clinical assessments, five studies used a screening tool [28,29,39,48,56], and four studies used patient self-reported dysphagia [49,52,54,55]. The included studies were conducted across fifteen countries, with studies most frequently conducted in Korea (n = 13), USA (n = 6), China (n = 3) and Japan (n = 3).

Outcome measures (Table 3). Many different outcome measures were used across the included studies targeting different domains within the area of OD. The most frequently used measures were the Penetration Aspiration Scale (PAS; 15 studies), the Functional Oral Intake Scale (FOIS; 8 studies), various water swallow tests (4 studies), and the Mann Assessment of Swallowing Ability (MASA; 3 studies). All other outcome measures were used in one or two studies only, confirming the substantial heterogeneity in outcome measures.

Interventions (Table 3). The included 37 studies comprised a range of behavioural interventions, delivered by various health professionals. The interventions were most frequently implemented by single allied health disciplines: occupational therapists in ten studies, speech pathologists in eight studies, physical therapists in two studies [36,48], and nursing staff in one study [55]. In five studies, more than one discipline was involved [23,27,28,33,48], and two studies reported caregivers as the intervention agent either as a single agent [24] or in addition to occupational therapists [22]. Nine studies did not specify disciplines involved in providing the interventions. The intervention dosage varied greatly, ranging from one training session [54] to exercise 3 times daily, 7 days per week for 42 days [25].

Behavioural intervention groups (Table 3). Of the 37 included studies, seven studies comprised three participant groups [19–21,23,25,26,38], whereas all other studies included two groups. Based on authors’ description of therapy contents, all intervention groups were categorised into compensatory, rehabilitative, and combined compensatory and rehabilitative interventions. Ten studies included different types of intervention groups (i.e., compensatory, rehabilitative and/or combined compensatory and rehabilitative intervention groups). Five studies included only compensatory groups [20,24,38,39,55], ten studies included only rehabilitative groups, and thirteen studies included only groups combining compensatory and rehabilitative interventions.

Most studies (n = 23) included a comparison group that received a type of dysphagia treatment often referred to as traditional therapy, standard swallow therapy, or conventional dysphagia treatment (CDT). Some studies also used the term usual care for CDT groups. CDT treatment could include counselling and the provision of information about swallowing and dysphagia, compensatory strategies (e.g., bolus modification and adjusted head positioning), rehabilitation, oromotor exercises and/or thermal stimulation. Three studies included a comparison group receiving medical standard care without dysphagia treatment [20,51,56]. In three studies, patients underwent sham dysphagia training [36,43,53]. Several studies compared two or three behavioural interventions without having a CDT or medical standard care group included [33,34,46,49,50,55].
3.3. Risk of Bias Assessment

The Begg and Mazumdar rank correlation procedure produced a tau of 0.305 (two-tailed \( p = 0.113 \)), indicating there is no evidence of publication bias. This meta-analysis incorporates data from 15 studies, which yield a \( z \)-value of 7.528 (two-tailed \( p < 0.001 \)). The fail-safe N is 207. This means that 207 ‘null’ studies need to be located and included for the combined two-tailed \( p \)-value to exceed 0.050. That is, there would need to be 13.8 missing studies for every observed study for the effect to be nullified. Both of these procedures (i.e., Begg and Mazumdar rank correlation and fail-safe N) indicate the absence of publication bias.

3.4. Methodological Quality

Risk of bias of the included RCTs was assessed using the RoB 2 tool. Figures 2 and 3 present the risk of bias summary per domain for individual studies and for all included studies. Most studies showed low risk of bias per domain, but more than half of the included studies (19/37) scored overall as having some concerns, with three studies identified as being at high risk.

![Figure 2. Risk of bias summary for all included studies (\( n = 37 \)) in accordance with RoB2.](image-url)
Figure 3. Risk of bias summary for individual studies (n = 37) in accordance with RoB2 [19–55].

Note. If one or more yellow or red circles (domains) have been identified for a particular study, the Overall score (last column) shows an exclamation mark, indicating that either the study shows some concerns (yellow circle with exclamation mark) or is at high risk (red circle with exclamation mark).
3.5. Meta—Analysis: Effect of Interventions

Twenty-one studies were included in the meta-analyses [21,22,24,25,28–31,34,35,40–46,49,51,52,54]. All study groups were categorised into compensatory interventions, rehabilitative interventions, combined compensatory and rehabilitative interventions, and no dysphagia intervention. Seventeen studies were excluded from meta-analyses: one study included patients with self-reported swallowing difficulties without confirmed OD diagnosis by instrumental assessment (VFSS or FEES) [48], four studies did not report on instrumental or clinical non-instrumental outcome data [20,28,37,40], ten studies provided insufficient data for meta-analysis [21,24,27,34,38,48,51,56,57], and two studies were excluded to reduce heterogeneity between studies [32,53].

Overall, within group analysis. (Figure 4). A significant, large pre-post intervention effect size was calculated using a random-effects model (z(35) = 8.047, p < 0.001, Hedges’ g = 1.139, and 95% CI = 0.862–1.416). Pre-post intervention effects varied greatly between studies, ranging from 0.058 to 5.732. Of the 36 intervention groups included in the meta-analysis, 19 groups showed large effect sizes (Hedges’ g > 0.8), six groups showed moderate effects sizes (0.5 < Hedges’ g ≤ 0.8), seven groups showed minor effect sizes (0.2 < Hedges’ g ≤ 0.5), and four groups showed negligible effect sizes (Hedges’ g ≤ 0.2). Between-study heterogeneity was significant (Q(35) = 152.938, and p < 0.001), with I² showing heterogeneity accounted for 77.115% of variation in effect sizes across studies.

| Study name                        | Subgroup within study | Statistics for each study | Hedges’s g and 95% CI |
|-----------------------------------|-----------------------|---------------------------|-----------------------|
| Carnaby et al. (2020a)            | Rehabilitation        | 1.052 0.359 0.129 0.349 1.756 2.932 0.003 |
| Carnaby et al. (2020b)            | Combination           | 0.784 0.353 0.125 0.52 1.476 2.219 0.028 |
| Choi et al. (2017a)               | Combination           | 2.057 0.430 0.185 1.214 2.899 4.783 0.000 |
| Choi et al. (2017b)               | Compensation          | 0.652 0.395 0.133 0.863 1.368 1.796 0.074 |
| Eom et al. (2017)                 | Rehabilitation        | 1.516 0.414 0.171 0.705 2.327 3.663 0.000 |
| Gao & Zhang (2017a)               | Rehabilitation        | 1.355 0.292 0.080 0.782 1.869 4.727 0.000 |
| Gao & Zhang (2017b)               | Rehabilitation        | 1.326 0.282 0.080 0.773 1.879 4.701 0.000 |
| Gao & Zhang (2017c)               | Rehabilitation        | 0.446 0.258 0.067 -0.060 0.552 1.727 0.084 |
| Hägglund et al. (2020a)           | Rehabilitation        | 0.058 0.326 0.106 -0.581 0.697 1.777 0.859 |
| Hägglund et al. (2020b)           | Rehabilitation        | 0.278 0.382 0.146 -0.471 1.026 0.727 0.467 |
| Heang et al. (2019a)              | Combination           | 0.464 0.416 0.173 -0.363 1.280 1.115 0.265 |
| Heang et al. (2019b)              | Combination           | 0.109 0.429 0.184 -0.731 0.949 0.255 0.799 |
| Jakobsen et al. (2019a)           | Rehabilitation        | 4.189 1.597 1.204 2.039 6.340 3.818 0.000 |
| Jakobsen et al. (2019b)           | Rehabilitation        | 0.635 0.589 0.346 -0.519 1.789 0.107 0.281 |
| Jang et al. (2019a)               | Combination           | 2.102 0.409 0.168 1.299 2.904 5.134 0.000 |
| Jang et al. (2019b)               | Combination           | 1.197 0.355 0.126 0.561 1.983 3.370 0.000 |
| Mepani et al. (2009a)             | Rehabilitation        | 0.378 0.528 0.290 -0.678 1.433 0.702 0.483 |
| Mepani et al. (2009b)             | Rehabilitation        | 1.077 0.620 0.384 -0.138 2.292 1.737 0.082 |
| Moon et al. (2018a)               | Combination           | 5.732 1.118 1.250 3.541 7.024 5.126 0.000 |
| Moon et al. (2018b)               | Combination           | 4.079 0.862 0.743 2.389 5.769 4.731 0.000 |
| Park et al. (2016)                | Rehabilitation        | 1.934 0.449 0.201 1.054 2.913 4.309 0.000 |
| Park et al. (2018a)               | Rehabilitation        | 1.690 0.483 0.233 0.743 2.638 3.500 0.000 |
| Park et al. (2018b)               | Rehabilitation        | 0.300 0.413 0.170 -0.509 1.169 0.726 0.468 |
| Park, Oh et al. (2019a)           | Rehabilitation        | 1.317 0.438 0.192 0.460 2.175 3.010 0.003 |
| Park, Oh et al. (2019b)           | Rehabilitation        | 0.647 0.405 0.164 -0.147 1.441 1.598 0.110 |
| Park, Lee et al. (2019a)          | Combination           | 1.574 0.365 0.133 0.858 2.290 4.308 0.000 |
| Park, Lee et al. (2019b)          | Combination           | 0.970 0.346 0.119 0.293 1.647 2.809 0.005 |
| Park et al. (2020a)               | Combination           | 2.364 0.468 0.219 1.446 3.282 5.047 0.000 |
| Park et al. (2020b)               | Combination           | 0.632 0.377 0.142 -0.106 1.370 1.687 0.063 |
| Steeke et al. (2016a)             | Rehabilitation        | 0.306 0.521 0.271 -0.715 1.327 0.587 0.557 |
| Steeke et al. (2016b)             | Rehabilitation        | 0.578 0.553 0.306 -0.506 1.663 1.045 0.296 |
| Tanamosale et al. (2019a)         | Combination           | 3.281 0.673 0.453 1.963 4.600 4.877 0.000 |
| Tanamosale et al. (2019b)         | Combination           | 2.767 0.612 0.575 1.587 3.967 4.519 0.000 |
| Tisch et al. (2017)               | Rehabilitation        | 0.351 0.257 0.086 -0.152 0.655 1.397 0.172 |
| Wolsert et al. (2020a)            | Compensation          | 0.148 0.274 0.075 -0.389 0.684 0.539 0.590 |
| Wolsert et al. (2020b)            | Compensation          | 0.146 0.255 0.065 -0.354 0.646 0.573 0.567 |
| Wolsert et al. (2020c)            | Compensation          | 1.139 0.142 0.020 0.862 1.416 8.047 0.000 |

Figure 4. Within intervention group pre-post meta-analysis [21,22,24,25,28–31,40–46,49,51,52,54,56]. Note. Refer to Table 2 for explanation of the subgroups.

Between subgroup analyses. Subgroup analyses (Table 4) were conducted comparing different types of interventions: behavioural interventions were compared with conventional dysphagia treatment (CDT), or no dysphagia therapy groups (Figure 5). Both behavioural interventions and CDT were categorised into mainly compensatory, rehabilitative, and combined compensatory and rehabilitative interventions. Overall, significant
treatment effects were identified favouring behavioural interventions. In particular, large effect sizes were found when comparing rehabilitative interventions with no CDT, and combined interventions with compensatory CDT. When comparing selected interventions based on commonalities across studies against CDT, significant, large effect sizes were found in favour of Shaker exercise, chin tuck against resistance exercise (CTAR), and expiratory muscle strength training (EMST). Most studies were conducted in stroke populations and showed significant, moderate effect sizes. Comparisons between outcome measures indicated at significant effects for PAS only.

Table 4. Between subgroup meta-analyses comparing intervention groups of included studies.

| Subgroup                                      | Hedge’s g | Lower Limit CI | Upper Limit CI | Z-Value | p Value |
|-----------------------------------------------|-----------|----------------|----------------|---------|---------|
| Intervention type                             |           |                |                |         |         |
| Combined vs. CDT (Combined) (n = 5)           | 0.610     | 0.263          | 0.957          | 3.446   | 0.001 * |
| Combined vs. CDT (Compensation) (n = 3)      | 1.180     | 0.362          | 1.998          | 2.828   | 0.005 * |
| Rehabilitation vs. CDT (Combined) (n = 1)    | 0.019     | -0.656         | 0.659          | 0.057   | 0.955   |
| Rehabilitation vs. CDT (Rehabilitation) (n = 3) | 0.178     | 0.304          | 1.133          | 3.395   | 0.001 * |
| Rehabilitation vs. No CDT (n = 3)            | 0.842     | 0.440          | 1.244          | 4.110   | <0.001 * |
| Selected interventions                        |           |                |                |         |         |
| Shaker vs. CDT (n = 2)                        | 1.038     | 0.300          | 1.776          | 2.756   | 0.006 * |
| CTAR vs. CDT (n = 3)                          | 1.045     | 0.427          | 1.663          | 3.316   | 0.001 * |
| EMST vs. no CDT (n = 2)                       | 0.819     | 0.389          | 1.250          | 3.733   | <0.001 * |
| Diagnostic groups                             |           |                |                |         |         |
| Acquired Brain Injury (n = 1)                 | 0.947     | -0.247         | 2.141          | 1.554   | 0.120   |
| Parkinson’s disease (n = 1)                   | 0.792     | 0.273          | 1.311          | 2.898   | 0.003 * |
| Stroke (n = 13)                                | 0.731     | 0.474          | 0.988          | 5.573   | <0.001 * |
| Outcome measures                              |           |                |                |         |         |
| Superior hyoid displacement (n = 1)           | 0.994     | -0.124         | 2.112          | 1.743   | 0.081   |
| MASA (n = 2)                                  | 0.512     | -0.574         | 1.599          | 0.925   | 0.355   |
| PAS (n = 11)                                  | 0.804     | 0.572          | 1.036          | 6.789   | <0.001 * |
| Tongue motility oromotor function (n = 1)     | 0.359     | -0.470         | 1.189          | 0.849   | 0.396   |

Notes. * Significant.

4. Discussion

Figure 5. Between subgroup meta-analysis for different types of interventions: behavioural interventions compared with conventional dysphagia treatment (CDT) or no dysphagia therapy [21,22,25,29–31,34,35,41–44,46,52]. Note. Refer to Table 2 for explanation of the subgroups.
This systematic review aimed to determine the effects of behavioural interventions in people with OD based on the highest level of evidence (RCTs) only. Findings from the literature were reported using PRISMA and meta-analysis procedures.

4.1. Systematic Review Findings

In total, 37 behavioural RCTs in OD were identified. Considering the high prevalence [3] and severe impact of OD on health [57], quality of life [5,58], and health-economics [59], the limited number of high-level evidence studies is concerning. RCTs are costly and usually require extensive funding [60]. Possibly, the general lack of awareness of OD [61] might place funding applications in this research area at a disadvantage when competing with well-known, life-threatening diseases such as cancer or stroke. Although OD is a symptom of these diseases, and many other underlying conditions, limited public knowledge persists, resulting in reduced understanding and recognition of the devastating consequences of OD, in both health-care and non-health-care practitioners [61].

Further, although RCTs are characterised by random allocation and allocation concealment, few of the included studies included sufficient reporting on the processes of randomization and blinding. These finding are in line with current literature on quality assessments of RCTs [62,63], confirming that the risk of selection bias [63] and the success of blinding methods in RCTs [62] can often not be ascertained due to frequent poor reporting.

When comparing behavioural RCTs in OD, several methodological challenges arise. Authors may use different definitions for OD or fail to provide sufficient details when reporting on the swallowing problems of the included patient populations. Also, several studies used non-instrumental assessments (i.e., patient self-report or a screening tool) to identify or confirm OD, making the comparison between studies precarious. The use of a screening tool is especially problematic in identifying OD and cannot act as confirmation of OD. A screening tool’s purpose is merely to identify patients at risk of OD, after which further assessment may confirm or refute the diagnosis [2]. Additionally, although instrumental assessment is considered the optimal tool for confirming OD diagnosis, VFSS and FEES protocols may differ (e.g., using different numbers of swallow trials, viscosities, and volumes).

Studies used a wide range of outcome measures to evaluate treatment effects. Since OD is a multidimensional phenomenon [64], different dimensions of OD may result in different therapy outcomes. For example, changes in dysphagia-related quality of life or oral intake do not necessarily correlate with findings from instrumental assessment. As such, to reduce heterogeneity in meta-analyses, patient self-report and oral intake measures were excluded. Also, some studies included outcome measures with poor or unknown psychometric properties, which in turn undermines the interpretation of treatment effects as data may not be valid or reliable. In addition, measures with weak responsiveness characteristics are not sensitive to treatment changes and should therefore be avoided as outcome measures aiming to determine intervention effects [2].

Most studies included a combined rehabilitative and compensatory intervention group or a rehabilitative intervention group, with only a few studies including exclusively compensatory groups. As the interventions classified as CDT comparison groups showed large variation as well, CDT comparison groups were categorised into similar group types (compensatory and/or rehabilitative CDT). Overall, terminology in the literature referring to CDT comparison groups was varied and complex. This was especially pertinent when interventions were not described in sufficient detail and descriptive terms such as “usual care” or “traditional therapy” did not provide further clarity on the type or content of CDT provided. Despite using categories to group different types of interventions, some degree of heterogeneity was inevitable. Interventions used different types of exercises or care, in distinct dosages, and were applied by different health care professionals. Therefore, it is challenging to identify the “active” ingredients of individual interventions, especially as most studies combined the use of different treatment strategies.
4.2. Meta-Analysis Findings

When considering meta-analyses for behavioural interventions, overall significant treatment effects were identified as favouring behavioural interventions over CDT and withholding dysphagia therapy. Most promising intervention approaches were rehabilitative interventions, which were associated with large effect sizes. Additionally, rehabilitative interventions such as Shaker exercise, CTAR exercise, and EMST showed significant, large effect sizes. However, since most studies included in the meta-analysis provided data on stroke patients only, future research still needs to confirm these findings in other diagnostic populations such as Parkinson’s disease, acquired brain injury or patients with head and neck oncology. As stated above, patient self-report and oral intake measures were excluded from meta-analyses to increase homogeneity between studies. Though self-report and oral intake data might be interesting for future meta-analyses, this would require additional RCTs to be published, as currently there is limited data available in the literature. Finally, future studies should report on treatment dosage and duration in more detail. Due to high heterogeneity between studies and incomplete reporting, no subgroup meta-analyses could be conducted for these variables.

4.3. Limitations

Although reporting of this review followed the PRISMA guidelines to reduce bias, some limitations are inherent to this study. As only RCTs published in English were included, some RCTs may have been excluded based on language criteria. In addition, meta-analyses were restricted because of heterogeneity of the included studies. As such, comparisons across studies are challenging and, generalisations and meta-analyses results should be interpreted with caution.

5. Conclusions

Meta-analyses for behavioural studies in oropharyngeal dysphagia identified an overall, significant, large pre-post interventions effect size. Significant treatment effects were identified favouring behavioural interventions over conventional dysphagia treatment. Notably, large effect sizes were found when comparing rehabilitative interventions with no dysphagia treatment and combined interventions with compensatory conventional dysphagia treatment. Selected interventions compared with conventional dysphagia treatment showed significant, large effect sizes in favour of Shaker exercise, CTAR, and EMST.

Behavioural interventions show promising effects in people with oropharyngeal dysphagia. Still, generalisations from this meta-analysis need to be interpreted with care due to high heterogeneity across studies.

Supplementary Materials: The following supporting information can be downloaded at: www.mdpi.com/article/10.3390/jcm11030685/s1, Table S1: PRISMA 2020 for Abstracts Checklist; Table S2: PRISMA 2020 Checklist.

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