Interventions for motor apraxia following stroke (Review)

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ABSTRACT

Background
Apraxia is a cognitive disorder that can occur after stroke. It prevents a person from carrying out a learned movement. Various interventions are used to treat apraxia but evidence of their benefit has been lacking.

Objectives
To determine which therapeutic interventions targeted at motor apraxia reduce disability.

Search strategy
We searched the Cochrane Stroke Group Trials Register (last searched November 2006). In addition, we searched the following databases: the Cochrane Central Register of Controlled Trials (CENTRAL) (The Cochrane Library Issue 3, 2006), MEDLINE (1966 to November 2007), EMBASE (1980 to November 2006), CINAHL (1982 to November 2006), PsycINFO (1974 to November 2006), the Research Index of the Occupational Therapy Journal (searched November 2006), REHABDATA (1956 to November 2006), the National Research Register (searched November 2006) and Current Controlled Trials Register (searched November 2006). We reviewed the reference lists of all articles that we identified as relevant. We made efforts to find both published and unpublished trials by writing to key authors and journals.

Selection criteria
Randomised controlled trials of therapeutic intervention for motor apraxia in stroke.

Data collection and analysis
One review author searched the titles, abstracts and keywords. Four review authors extracted data and analysed trial quality. We contacted investigators for further details of trials if necessary.

Main results
Three trials including a total of 132 participants were included in the review. There was evidence of a small and short-lived therapeutic effect in the two studies that reported change in activities of daily living (102 participants) but this was not considered clinically significant and did not persist at the longer-term follow up.

Authors’ conclusions
There is insufficient evidence to support or refute the effectiveness of specific therapeutic interventions for motor apraxia after stroke. Further research of higher quality is required. As we did not review whether patients with apraxia benefit from rehabilitation input in general, they should continue to receive general stroke rehabilitation services.

PLAIN LANGUAGE SUMMARY
Interventions for motor apraxia following stroke

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People with motor apraxia after stroke often have difficulty carrying out everyday activities such as making a hot drink. Some people cannot select the right object at the right time or have difficulty using objects (such as a spoon) correctly. Apraxia is not due to muscle weakness or sensory loss. Instead it seems to be a loss or disturbance of the conceptual ability to organise actions to achieve a goal. This review of three studies, including 132 participants, suggests that further high quality research is required before specific treatment techniques can be accepted or rejected. Patients with apraxia should continue to receive general stroke rehabilitation services but better quality research is needed to identify optimal apraxia treatments.

**BACKGROUND**

The World Health Organization has defined stroke as ‘a syndrome of rapidly developing clinical signs of focal (or global) disturbance of cerebral function, with symptoms lasting 24 hours or longer leading to death, with no apparent cause other than vascular in origin’ (WHO 1978). Stroke is the largest disabling condition in England and Wales with 100,000 first strokes occurring each year (Blais 1994). Stroke can affect people’s physical, sensory and cognitive abilities (Wade 1985). The Stroke Association estimates that in the UK 300,000 of the 60 million population are living with disabilities caused by a stroke (Westcott 2000).

Apraxia is a neuropsychological deficit that can affect stroke patients. It refers to ‘disorders of the execution of learned movement which cannot be accounted for by either weakness, inco-ordination, or sensory loss, or by incomprehension of or inattention to command’ (Geschwind 1975). In this review we shall confine the discussion of apraxia to that affecting the limbs. Apraxia of speech is dealt with in a separate Cochrane review (West 2005).

Motor apraxia is difficult to diagnose. The available tests are inconsistent and appear to test for different aspects of apraxia (Butler 2002). The taxonomy of motor apraxia has been disputed, but many clinicians and researchers now support the classical idea that there are two forms: ideomotor and ideational (Liepmann 1920). Others have described motor apraxia in functional terms, for example dressing apraxia and the apraxia of gait. These classifications have been disputed as they describe the affected functional task rather than the underlying condition (Geschwind 1985). Ideomotor apraxia can affect the patient by hindering their ability to select, sequence and use objects (Heilman 1985) and it is thought to affect people more in test situations than in normal activities of daily living (ADL). Patients with ideational apraxia are unable to perform a skilled activity because they have lost the conceptual ability to organise the actions required to achieve their goal (Jackson 1999). For example, they may attempt to put clothes on the wrong part of their body. There does not, however, appear to be a clear consensus on the definitions of ideomotor and ideational apraxia (Tate 1995).

The reported prevalence of motor apraxia after stroke is inconsistent. There is evidence to suggest that apraxia affects both left and right-brain damaged patients, with it being more prevalent in the left (Rotthi 1997). Both the anterior and posterior lesions in the left hemisphere are known to produce apraxic symptoms, as this is the dominant hemisphere for the storage and execution of learned movements (Kareken 1998). Original studies showed that 50% of patients with right-sided hemiplegia suffered from motor apraxia (Liepmann 1905). This has been confirmed by another study (De Renzi 1980).

Apraxia is thought to have an adverse influence on ADL independence (Goldenberg 1998; Sundet 1988). Research into the different therapeutic interventions available to treat apraxia is limited. Types of interventions include:

- **strategy training in daily living activities:** this technique teaches internal (for example, the patient is taught to verbalise and implement the task steps at the same time) or external (for example, when aids are used to overcome a functional barrier) compensatory strategies that enable a functional task to be completed. These strategies will not have been used prior to the stroke (Van Heugten 1998);
- **sensory stimulation:** stimulations including deep pressure, sharp and soft touch are applied to the patients’ limbs (Butler 1994);
- **propiroceptive stimulation:** the patient leans on and puts weight through their upper and lower limbs;
- **cuing, verbal or physical prompts:** given to enable each stage of the task to be completed;
- **chaining (forward or backward):** the task is broken down into its component parts. Using backward chaining the task is completed with facilitation from the therapist apart from the final component, which the patient carries out unaided. If successful next time further steps are introduced. Forward chaining is the reverse of backward chaining;
- **normal movement approaches:** the therapist facilitates the body through normal movement patterns.

Rehabilitation can occur at any phase post stroke. There is a conceptual distinction between the effects a disease may have at different levels (WHO 2001): impairment, activity (disability) and participation (handicap). Therapists’ provision of aids and environmental adaptations aims to help the person adapt to their impairment rather than change the underlying impairment itself. Some rehabilitation approaches may be aimed at the level of impairment.
The task of this review is to systematically consider the evidence from randomised controlled trials on the effectiveness of therapeutic interventions aimed specifically at altering motor apraxia following stroke.

**OBJECTIVES**

The main questions we wish to address are as follows.

1. In stroke patients with motor apraxia who are undergoing rehabilitation, do therapy interventions targeted at motor apraxia achieve a sustained reduction in disability compared with no or placebo intervention six months after treatment?

2. In this population, is one specific targeted intervention (compared with another specific targeted intervention) more likely to achieve a sustained reduction in disability?

**CRITERIA FOR CONSIDERING STUDIES FOR THIS REVIEW**

**Types of studies**

We included randomised controlled trials of interventions for stroke patients with motor apraxia. We would have excluded from analysis second and subsequent phases of cross-over trials, as the design would not be appropriate in this context.

**Types of participants**

The review was confined to data from reports of studies on adult patients with motor apraxia (irrespective of the definition of apraxia used by the authors of the study) following a stroke. We excluded trials that included participants whose deficits were the result of head trauma, brain tumour, or other brain damage unless a subgroup of stroke patients could be identified for whom there were separate results, or more than 75% of patients in the sample are stroke patients. All types of apraxia (that is ideomotor and ideational) were considered for inclusion except apraxia of speech and oral apraxia. Apraxia of speech has been covered in a separate Cochrane review (West 2005).

**Types of intervention**

We included trials in which a comparison was made between an 'active' treatment group that received one of the various motor apraxia interventions and a control group that received either an alternative motor apraxia intervention, placebo or none. Possible treatment interventions included: tactile and proprioceptive stimulation, strategy training in daily living activities, cueing, chaining, (forward or backward) and normal movement approaches. We excluded trials involving only drug therapies. We recorded duration and quantity of intervention.

**Types of outcome measures**

The primary outcome was the average level of independence in activities of daily living, as defined by the original authors, at six months after therapy. Recognised measures, for example the Barthel Index (Mahoney 1965), the Assessment of Motor and Process Skills (Fisher 1994) and the Functional Independence Measure (Keith 1987) were included.

Secondary outcomes included:

1. Independence in ADL at the scheduled end of the intervention (ordinal);
2. Independence in ADL at 12 months (ordinal);
3. Death (binary);
4. Quality of life measures (ordinal);
5. Ability to gesture/pantomime/use objects (ordinal);
6. Effects on family and carer, e.g. Carer Strain Index, measures of carer’s mood (ordinal);
7. Carer and family perceptions of outcome (ordinal);
8. Economic resources (continuous);
9. Apraxic patient’s mood (ordinal);
10. Adverse events, e.g. fatigue, falls, accident rates (binary).

**SEARCH METHODS FOR IDENTIFICATION OF STUDIES**

See: Cochrane Stroke Group methods used in reviews.

(1) We searched the Cochrane Stroke Group Trials Register, which was last searched by the Review Group Co-ordinator in November 2006. In addition, we searched the Cochrane Central Register of Controlled Trials (CENTRAL) (The Cochrane Library Issue 3, 2006), MEDLINE (1966 to November 2006), EMBASE (1980 to November 2006), CINAHL (1982 to November 2006), PsycINFO (1974 to November 2006), the Research Index of the Occupational Therapy Journal (searched November 2006), REHABDATA (1956 to November 2006), the National Research Register (searched November 2006) and Current Controlled Trials Register (searched November 2006).

The search strategy for MEDLINE is given below and this was modified for the other databases.

**Database MEDLINE (Ovid) 1966 to November 2006**

1 exp cerebrovascular disorders/
2 (stroke$ or poststroke$ or cva$).tw.
3 (cerebrovascular$ or cerebral vascular).tw.
4 (cerebral or cerebellar or brainstem or vertebrobasilar).tw.
5 (infarct$ or isch?emi$ or thrombo$ or apoplexy or emboli$).tw.
6 4 and 5
7 (cerebral or intracerebral or intracranial or parenchymal).tw.
8 (brain or intraventricular or brainstem or cerebellar).tw.
9 (infratentorial or supratentorial or subarachnoid).tw.
10 7 or 8 or 9
11 (haemorrhage or hemorrhage or haematoma or hematoma).tw.
12 (bleeding or aneurysm).tw.
13 11 or 12
14 10 and 13
We had planned to handsearch a number of relevant journals. However, after checking the Master List of journals being searched by The Cochrane Collaboration to avoid duplication of effort (http://www.cochrane.us/masterlist.asp), we found that the selected journals had already been handsearched. The resulting trials would therefore be found from our search of the Cochrane Central Register of Controlled Trials.

(3) We searched the reference lists of all relevant references.

(4) In order to identify further published and unpublished trials we contacted authors of published apraxia articles and wrote to appropriate journals (Clinical Rehabilitation, British Journal of Occupational Therapy, Physiotherapy Frontline and The Psychologist).

METHODS OF THE REVIEW

Selection of trials
One review author (CW) searched titles, abstracts and keywords of both published and unpublished papers to assess their eligibility for inclusion using a systematic approach. Only papers that obviously did not meet the eligibility criteria were discarded. Articles that may have met the inclusion criteria were obtained in full and screened by CW. All review authors read the remaining studies and formed a consensus on the final inclusion and data extraction.

Quality assessment
We described the methodological quality of the included studies for the following aspects:

- concealment of allocation (whether adequate, inadequate, or unclear);
- type of design (e.g. parallel, factorial, cross-over);
- blinding to allocation (of therapist, patient and outcome assessment);
- definition of terms (e.g. of stroke, apraxia, outcome, and intervention);
- intention-to-treat analysis (whether undertaken, possible from report, impossible or unclear);
- completeness of follow up (proportion of randomised patients in analysis).

Data extraction
In addition to outcome data the following were documented by CW and one other review author: (1) settings (e.g. hospital, community, nursing home); (2) type of intervention; (3) length of rehabilitation; (4) profession(s) involved; (5) co-interventions implemented; (6) length of disease; (7) level of severity; (8) presence of other symptoms that may affect the level of disability (e.g. hemiplegia, unilateral spatial neglect); and (9) tools the authors used to identify motor apraxia. We requested information that was unclear or missing from the reports from the corresponding author.

Data analysis
Our primary analysis pooled all therapeutic studies of active intervention versus no or placebo treatment to address objective (1) above. To address objective (2), we also analysed subgroups of studies categorised according to therapeutic approach, as outlined under 'Types of interventions'. This included a comparison of each approach versus no or placebo treatment, and will include direct comparisons of different approaches if any are identified in future updates of this review.
We have treated activities of daily living (ADL) and other ordinal scales for the secondary outcomes as continuous outcomes unless and until accepted meta-analytic techniques for ordinal outcome data become available. We abstracted, calculated or requested means and standard deviations. For all binary outcomes, we incorporated deaths in the worse outcome category. For practical reasons, we excluded deaths from outcomes that were treated as continuous. Death rates between the two groups were low and similar because studies only included patients who were well enough to undergo rehabilitation for motor apraxia. Any imbalance in death rates between the groups in future updates will be discussed, including descriptive consideration of whether analyses of raw data from individual trials could alter conclusions.

Our intention was to extract mean (SD) for the primary outcome, and this was possible for included studies. If this is not the case in future updates, we will extract and compare binary data for the primary outcome as an additional secondary analysis.

We combined results for continuous outcomes using weighted mean difference by a fixed-effect model. However, it is anticipated that future studies may use different scales to measure the same underlying constructs. If this is the case, we will use the standardised mean difference and results translated back into one of the original scales for reporting purposes. We combined results for binary outcomes using the Feto-modified odds ratio (OR), and translated these to risk differences across the observed range of control group rates for reporting purposes. We noted and discussed statistical heterogeneity.

We carried out sensitivity analyses on the primary outcome. These included use of a random-effects analysis, omission of studies that do not describe an adequate method of allocation concealment, and imputing values for missing data if appropriate.

**DESCRIPTION OF STUDIES**

There were no excluded studies as no studies that appeared to meet the eligibility criteria were found not to on closer examination. Data from 132 participants in three studies were included (Donkervoort 2001; Edmans 2000; Smania 2000). Smania 2000 reported data for 13 patients but we have only included data for the first 10 patients who were appropriately randomised (see ‘Methodological quality of included studies’). Edmans 2000 provided segregated data on the nine patients with apraxia included in her published report.

The participants all had lesions in the left hemisphere. Apraxia was defined in Donkervoort 2001 using the De Renzi test (De Renzi 1980), in Smania 2000 using the Van Heugten test (Van Heugten 1999), and in Edmans 2000 using the test by Kertesz and Ferro (Kertesz 1984). The mean ages of groups were between 63 and 70 years. The sex (male/female) of the experimental groups was 64/49 (Donkervoort 2001), 8/2 (Smania 2000) and 3/6 (Edmans 2000). The study participants came from the Netherlands (Donkervoort 2001), Italy (Smania 2000), and England (Edmans 2000) and were from either a rehabilitation unit (Donkervoort 2001; Edmans 2000; Smania 2000) or nursing home (Donkervoort 2001). The time since stroke was a mean of about 100 days (Donkervoort 2001), and ranged from two to 36 months (Smania 2000) and from 22 to 76 days (Edmans 2000). In the Donkervoort study (Donkervoort 2001) 56 (19%) participants had recurrent stroke, but none had a history of apraxia prior to their current stroke. There was no previous history of cerebrovascular attacks in the stroke patients participating in the Smania study (Smania 2000), and status was not reported in the Edmans study (Edmans 2000). All studies excluded people with marked psychiatric problems.

The comparisons in the studies differed. Donkervoort 2001 used strategy training (integrated into usual occupational therapy) compared to usual occupational therapy. Smania 2000 compared gesture training for apraxia with conventional treatment for aphasia (Smania 2000). Edmans 2000 compared two specific methods for apraxia in addition to standard occupational therapy: transfer of training and functional approach. As the latter is more standard practice, we have chosen arbitrarily to treat this as the control group. Donkervoort 2001 reported that the experimental group had on average 25 occupational therapy sessions lasting in total 15 hours whilst the control group had 27 occupational therapy sessions with a total of 19 hours, during an eight week period. Patients in Smania 2000 received training sessions of approximately 50 minutes duration three times a week. The gesture training stopped once all training sections were completed, or a maximum of 35 treatment sessions (approximately 11 weeks). In Edmans 2000, participants received training for 2.5 hours per week for six weeks.

In Donkervoort 2001, the assessment of apraxia was made by a trained researcher following clinical screening by the medical team. The intervention was delivered by occupational therapists and assessment made by a blinded research assistant. The professions involved in assessment of eligibility, intervention and outcome assessment are not clear in Smania 2000. In Edmans 2000, a psychologist assessed apraxia at the outset, occupational therapists delivered the interventions, and outcomes were assessed both by nurses and an independent, blinded occupational therapist.

The outcomes used in the studies were different. Donkervoort 2001 reported as primary outcome the Van Heugten (Van Heugten 1999) measure of ADL at end of intervention and at five months after initial assessment, but also reported Barthel among secondary outcomes. Smania 2000 reported a number of impairment outcomes at the end of intervention, but nothing regarding activities of daily living. Edmans 2000 reported a number of outcomes including the Barthel measured both by nurses and occupational therapists at the end of intervention. We have used the occupational therapist assessments in the analyses.
**METHODOLOGICAL QUALITY**

All included studies claimed to be randomised controlled trials using two-group parallel designs. Standard, though different, assessments of apraxia and outcomes were used. Due to the nature of the interventions it would not have been possible to blind therapists or patients.

Donkervoort 2001 randomised participants using sequentially numbered, non-transparent, sealed envelopes prepared from random number tables. Allocation was stratified by institution type, time since stroke and apraxia score, and a Zelen correction (Zelen 1974) was used to ensure balance. The outcome assessments were carried out by a blinded research assistant. Patients were not specifically informed which intervention they were receiving, although clearly the interventions would not have appeared similar. Stroke was defined using the WHO criteria (WHO 1989). The trialists referred to an article in which the intervention was defined in sufficient detail to replicate (Van Heugten 1998). Of 113 randomised patients, 108 (96%) underwent baseline assessment, 97 (86%) were assessed at the end of intervention, and 86 (76%) at the final assessment. Reasons for withdrawal at each stage were reported and balanced between the groups. Analyses were by intention to treat for those patients with outcome data.

Smania 2000 used simple randomisation on the first 10 subjects without mention of concealment. After noticing an imbalance the following three subjects were assigned to the control group and their data have been excluded from our analyses. There was no mention of blinding of outcome assessment, which is a potential source of avoidable bias. Stroke was defined by computerised tomography (CT) scan and clinical evidence of left-sided, unilateral vascular lesions. The intervention was defined in sufficient detail to replicate. There were complete follow-up data for the 10 included patients.

Edmans 2000 described a randomisation scheme using pre-prepared envelopes from random number tables. Edmans informed the review authors that allocations were stored in sealed, opaque, numbered envelopes, only opened at the time of recruitment in the presence of a witness. The outcome assessments were carried out independently by a blinded nurse and occupational therapist. The post-treatment assessor was blinded to allocation. No definition of stroke was given. Intervention details were not provided in the study or a later paper. Some randomised patients were not assessed for apraxia due to language impairment. Complete follow-up data were made available to this review for the nine patients assessed to have apraxia.

**RESULTS**

The graphs of continuous outcomes are set so that values to the right favour the experimental group. For binary outcomes, lower odds in the experimental group are always shown to the left. For adverse outcomes (such as death) this means that values to the left favour the experimental group.

Our protocol specified comparison of the average levels of independence in activities of daily living. Presented below are comparisons of the average changes from baseline in these levels. These change score analyses have been chosen because they usually provide more precise estimates of the same treatment effects in the randomised trial setting.

**Comparison 01.01: Change in Barthel at six months after end of therapy**

Only Donkervoort 2001 reported on the primary outcome described in this review’s protocol. Using the Barthel ADL Index, the study did not find evidence of a lasting difference in functional performance six months post stroke: mean difference (MD) 0.17, 95% confidence interval (CI) -1.41 to 1.75, P = 0.83, in favour of the experimental group.

**Comparison 01.02: Change in Barthel at end of therapy**

Donkervoort 2001 and Edmans 2000 both reported the Barthel at end of intervention, and reported very similar group differences. The overall MD was 1.28, 95% CI 0.19 to 2.38, P = 0.02, in favour of the experimental group.

**Comparison 01.03: Change in Barthel at 12 months after end of therapy**

No trials reported data for this outcome.

**Comparison 01.04: Death**

There were no deaths in the studies of Edmans 2000 or Smania 2000, but seven in the study by Donkervoort 2001: odds ratio (OR) 0.41, 95% CI 0.09 to 1.9, P = 0.25, in favour of the experimental group but providing no evidence of differential death rates.

**Comparison 01.05: Quality of life measures**

No trials reported data for this outcome.

**Comparison 01.06: Ability to gesture, pantomime, use real objects**

Only Smania 2000 reported on this outcome, using both ability to gesture and to use real objects: MD for gesture training 8.4, 95% CI -15.8 to 32.6 points on a 0 to 72 scale, P = 0.50 in favour of the experimental group. MD for using real objects 1.2, 95% CI -3.2 to 5.6 points on a 0 to 14 scale, P = 0.59, in favour of the experimental group but again providing no evidence of differential ability.

**Comparison 01.07: Effects on family and carer**

No trials reported data for this outcome.

**Comparison 01.08: Carer and family perceptions**

No trials reported data for this outcome.

**Comparison 01.09: Economic resources**

No trials reported data for this outcome.
Comparison 01.10: Apraxic patient’s mood
No trials reported data for this outcome.

Comparison 01.11: Adverse events
No trials reported data for this outcome.

DISCUSSION

Only Donkervoort 2001 reported on the primary outcome for this review. Using the Barthel Index the study did not find evidence of a lasting difference in functional performance six months post stroke. This review does however suggest that therapeutic intervention produces a small but statistically significant improvement on the Barthel immediately after intervention as both Donkervoort 2001 and Edmans 2000 found in favour of the experimental group. These results whilst encouraging have limited application for clinical practice due to the small effect and the fact that it did not persist at follow up. No studies compared one intervention with any other. Only Smania 2000 reported on test performance, for example the ability to gesture and the use of objects. Neither was statistically significant. Death rates were low and similar for all the studies. This was expected as only patients that were well enough to undergo rehabilitation would have been included. No studies reported on quality of life measures, effects on family and carer, their perceptions of outcome, economic resources, mood or adverse events. If future research is carried out it would be appropriate for these to be used as secondary outcome measures.

The review found and included only three trials with a small number of participants (132). All the trials used different therapeutic interventions, including strategy training (Donkervoort 2001), a transfer of training approach (that is, practising one task with the aim of it generalising to related tasks) (Edmans 2000), and gesture training (Smania 2000). Not all the therapeutic interventions suggested in the literature have been evaluated. The quantity of treatment intervention varied between 15 hours and 29 hours and duration was from six weeks to 19 weeks. The assessment tools used to diagnose apraxia were all different and we are unsure whether they actually measure the same underlying construct. The participants came from rehabilitation units (Donkervoort 2001; Edmans 2000; Smania 2000) and nursing homes (Donkervoort 2001). It is not clear whether participants from rehabilitation units in England and Italy and nursing homes in the Netherlands are comparable in terms of level of dependency. The interventions were only reported in enough detail to replicate in two of the three studies. Edmans 2000 is to report on the intervention in a future article. Without detail of the intervention a trial is of little clinical value.

Donkervoort 2001 used adequately concealed randomisation utilising sequentially-numbered, non-transparent, sealed envelopes, prepared from random number tables. Edmans 2000 used a similar process but the recruiter prepared the envelopes prior to allocation. This is a potential source of bias. It would be preferable if the recruiter were not involved in the preparation of the envelopes. Smania 2000 reported using simple randomisation on the first 10 patients, but once an imbalance was noticed a ‘restricted randomisation scheme’ was implemented without mention of concealment. The randomisation process is unclear. Donkervoort 2001 and Edmans 2000 reported using a blinded outcome assessor whilst Smania 2000 did not mention blinding. This is a possible source of bias.

In summary, the review has not found strong evidence to support therapeutic intervention for motor apraxia in stroke patients. We have found no evidence that the impairment of motor apraxia is altered, or that intervention aimed specifically at motor apraxia alters disability. This should not be misinterpreted as evidence that rehabilitation does not work for patients with motor apraxia.

The quality of the studies is acceptable for the review but there are study limitations as outlined above. The findings of this review suggest that good quality randomised controlled trials are warranted. Apraxic assessments used in future studies need to measure both the level of impairment and activity (WHO 2001). Impairment measures are useful for describing the sample and the type and severity of motor apraxia. This is needed for decisions about whether results from the samples studied can be generalised to a typical heterogeneous clinical population. It is also important for future researchers to consider evaluating their treatment in terms of the patients’ opinion of outcome.

AUTHORS’ CONCLUSIONS

Implications for practice
Specific therapeutic intervention for motor apraxia following stroke cannot be supported or refuted by results from randomised controlled trials.

Implications for research
There is a need for more and higher quality trials of therapeutic intervention for motor apraxia. Trials should be sufficiently large to detect functionally meaningful differences in long-term outcome. Interventions should be explicitly defined and outcome measures need to include how apraxia affects everyday life.

POTENTIAL CONFLICT OF INTEREST
None known.

ACKNOWLEDGEMENTS

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**SOURCES OF SUPPORT**

External sources of support

- North West NHS R&D Executive UK

Internal sources of support

- No sources of support supplied

**REFERENCES**

References to studies included in this review

Donkervoort 2001 [published data only]


Edmans 2000 [published and unpublished data]


Smania 2000 [published data only]


Additional references

Blais 1994


Butler 1994


Butler 2002


De Renzi 1980


Fisher 1994


Geschwind 1975


Geschwind 1985


Goldenberg 1998


Heilman 1985


Jackson 1999


Kareken 1998


Keith 1987


Kertesz 1984


Liebermann 1920


Liebermann 1905


Mahoney 1965

Characteristics of included studies

<table>
<thead>
<tr>
<th>Study</th>
<th>Donkervoort 2001</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methods</td>
<td>A randomised, single blind, controlled trial design. Patients were randomised using sealed envelopes prepared from random number tables. Patients were pre-stratified on institution type, time since stroke and apraxia score and a Zelen correction was used to prevent unequal distribution.</td>
</tr>
<tr>
<td>Participants</td>
<td>Netherlands</td>
</tr>
<tr>
<td>113 left stroke</td>
<td></td>
</tr>
<tr>
<td>Exptl n=56, cntrl=57</td>
<td></td>
</tr>
<tr>
<td>Mean age: exptl 68, cntrl 63</td>
<td></td>
</tr>
<tr>
<td>Sex (male/female): exptl 29/27, cntrl 35/22</td>
<td></td>
</tr>
<tr>
<td>Inclusion criteria:</td>
<td>left hemisphere stroke, apraxia, staying on an inpatient unit (15 rehabilitation centres and 35 nursing homes)</td>
</tr>
<tr>
<td>Exclusion:</td>
<td>history of apraxia, stroke has occurred less than 4 weeks or more than 2 years ago, age younger than 25 years and older than 95 years, history of traumatic brain damage, brain tumour, psychiatric history</td>
</tr>
<tr>
<td>Professional assessing apraxia at onset was a trained researcher following screening by the medical team</td>
<td></td>
</tr>
<tr>
<td>Interventions</td>
<td>Strategy training (integrated into usual occupational therapy) compared to occupational therapy</td>
</tr>
<tr>
<td>Strategy training:</td>
<td>teaching the patient internal/external compensatory approaches to assist ADL performance</td>
</tr>
<tr>
<td>Intervention period 8 weeks</td>
<td></td>
</tr>
<tr>
<td>Intervention was delivered by occupational therapists</td>
<td></td>
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<tr>
<td>The intervention was defined in enough detail in a further study (see 'Additional references', Van Heugten 1999)</td>
<td></td>
</tr>
</tbody>
</table>

* Indicates the major publication for the study
Characteristics of included studies (Continued)

Outcomes
Outcomes were measured at baseline, 8 weeks and 5 months
Outcomes collected: ADL measures (Van Heugten measure of ADL, Barthel ADL Index, ADL judgement list filled in independently by the OT and patient)
Apraxia, motor functioning (Motricity Index, functional motor test), additional tests (verbal comprehension, memory, neglect, mental status)
Assessment was made by a blinded research assistant

Notes
Allocation by random number table
Blocks of size 2 plus Zelen correction could make allocation predictable

Allocation concealment
A – Adequate

Study
Edmans 2000

Methods
Randomised, single blinded controlled trial. Used a randomisation scheme using pre-prepared envelopes from random number tables. Edmans informed the review authors that the recruiter prepared the allocations prior to the study. Allocations were stored in sealed, opaque, numbered envelopes, only opened at the time of recruitment in the presence of a witness

Participants
UK Nottingham Stroke Unit
80 left and right hemiplegic participants in trial, data from 9 apraxics were abstracted. 6 apraxics in the functional approach (mean age 70) and 3 in the transfer of training approach (mean age 69). All left hemisphere strokes
Inclusion criteria: all ages, able to complete the RPAB, functional use in one hand, patient or family able to give consent
A psychologist assessed for the apraxia at the outset

Interventions
Comparison of the transfer of training and functional treatment approaches
Transfer of training: practising one perceptual task will affect the performance on other perceptual tasks, i.e. the cause of the perceptual problem is treated
The functional approach: repetitive practice of specific daily living tasks. Intervention given for 2.5 hours per week for 6 weeks in additional to general OT
OTs delivered the interventions

Outcomes
The Barthel Index, Edmans ADL Index and RPAB assessments were completed before and immediately post intervention
Other routine assessments were also collated, e.g. the apraxia test by Kertesz and Ferro
Outcomes were assessed by nurses and an independent, blinded OT
Intervention was to be published by a later article

Notes
Patients transferred to the stroke unit were participating in an evaluation study, the selection criteria included: medically stable, transfer with 2 nurses, no discharge date, able to tolerate 30 minutes of treatment, able to complete 2 out of 4 specified functional tasks

Allocation concealment
C – Inadequate

Study
Smania 2000

Methods
Randomised, controlled trial
First 10 patients assigned to expvl/cntrl group
Following 3 used a restricted randomisation scheme placed in cntrl group; the last 3 were not included in this review

Participants
Italy Neurological Rehabilitation Unit
10 strokes accepted into the review: expvl 6, cntrl 4
Mean age: expvl 69.3 years, cntrl 63 years
Sex (male/female): expvl 5/1, cntrl 3/1
Duration of stroke: expvl mean 14.7 months, cntrl mean 18 months
Neurologic severity (range 0-18): expvl mean 6.5, cntrl mean 7.5
Inclusion criteria: limb apraxia, length of illness at least 2 months, right handed, left hemisphere stroke
Exclusion: history of cerebrovascular attacks or psychiatric disorders
Professional assessing eligibility was not clear

Interventions
Exptl: gesture training for apraxia
Cntrl: conventional treatment for aphasia.
The experimental group program consisted of gesture production exercises, 35 intervention sessions, each lasting 50 minutes or a maximum of 35 treatment sessions
Professional assessing intervention was not clear

Outcomes
A battery of tests including an oral apraxia test, a constructional apraxia test and 3 limb praxic function tests.
No tests regarding ADL were carried out
Professional assessing outcome was not clear
The intervention was clear enough to replicate

Notes
Only the first 10 assigned have been included in the study as they were truly randomised
Large difference in stroke duration between exptl and cntrl groups

Allocation concealment
B – Unclear
ADL: activities of daily living
cntrl: control
exptl: experimental
OT: occupational therapy/therapist
RPAB: Rivermead Perceptual Assessment Battery

ANALYSES

Comparison 01. Experimental therapy versus standard care

<table>
<thead>
<tr>
<th>Outcome title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
<th>Effect size</th>
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<tbody>
<tr>
<td>01 Change in Barthel at six months after end of therapy</td>
<td>1</td>
<td>83</td>
<td>Weighted Mean Difference (Fixed) 95% CI</td>
<td>0.17 [-1.41, 1.75]</td>
</tr>
<tr>
<td>02 Change in Barthel at end of therapy</td>
<td>2</td>
<td>102</td>
<td>Weighted Mean Difference (Fixed) 95% CI</td>
<td>1.28 [0.19, 2.38]</td>
</tr>
<tr>
<td>03 Change in Barthel at 12 months after end of therapy</td>
<td>0</td>
<td>0</td>
<td>Weighted Mean Difference (Fixed) 95% CI</td>
<td>Not estimable</td>
</tr>
<tr>
<td>04 Death</td>
<td>3</td>
<td>132</td>
<td>Peto Odds Ratio 95% CI</td>
<td>0.41 [0.09, 1.89]</td>
</tr>
<tr>
<td>05 Quality of life measures</td>
<td>0</td>
<td>0</td>
<td>Weighted Mean Difference (Fixed) 95% CI</td>
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<tr>
<td>06 Ability to gesture, pantomime, use real objects</td>
<td>0</td>
<td>0</td>
<td>Weighted Mean Difference (Fixed) 95% CI</td>
<td>Subtotals only</td>
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<tr>
<td>07 Effects on family and carer</td>
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<td>0</td>
<td>Weighted Mean Difference (Fixed) 95% CI</td>
<td>Not estimable</td>
</tr>
<tr>
<td>08 Carer and family perceptions</td>
<td>0</td>
<td>0</td>
<td>Weighted Mean Difference (Fixed) 95% CI</td>
<td>Not estimable</td>
</tr>
<tr>
<td>09 Economic resources</td>
<td>0</td>
<td>0</td>
<td>Weighted Mean Difference (Fixed) 95% CI</td>
<td>Not estimable</td>
</tr>
<tr>
<td>10 Apraxic patient's mood</td>
<td>0</td>
<td>0</td>
<td>Weighted Mean Difference (Fixed) 95% CI</td>
<td>Not estimable</td>
</tr>
<tr>
<td>11 Adverse events</td>
<td>0</td>
<td>0</td>
<td>Odds Ratio (Fixed) 95% CI</td>
<td>Not estimable</td>
</tr>
</tbody>
</table>

COVER SHEET

Title
Interventions for motor apraxia following stroke

Authors
West C, Bowen A, Hesketh A, Vail A

Contribution of author(s)
Carolyn West, Audrey Bowen and Andy Vail obtained funding for the production of this review from the North West Region NHS Executive under their Research Development Fund scheme.
Carolyn West wrote the protocol and review with the assistance of Andy Vail, Audrey Bowen and Anne Hesketh.
Carolyn West is an occupational therapist, Audrey Bowen is a psychologist, Andy Vail is a medical statistician and Anne Hesketh is a speech and language therapist.

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Review first published 2008/1

Date of most recent amendment 25 September 2007

Date of most recent SUBSTANTIVE amendment 25 September 2007

What's New Information not supplied by author

Date new studies sought but none found Information not supplied by author

Date new studies found but not yet included/excluded Information not supplied by author

Date new studies found and included/excluded Information not supplied by author

Date authors' conclusions section amended Information not supplied by author

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Cochrane Library number CD004132

Editorial group Cochrane Stroke Group

Editorial group code HM-STROKE
### Analysis 01.01. Comparison 01 Experimental therapy versus standard care, Outcome 01 Change in Barthel at six months after end of therapy

Review: Interventions for motor apraxia following stroke  
Comparison: 01 Experimental therapy versus standard care  
Outcome: 01 Change in Barthel at six months after end of therapy

<table>
<thead>
<tr>
<th>Study</th>
<th>Experimental</th>
<th>Control</th>
<th>Weighted Mean Difference (Fixed)</th>
<th>Weight (%)</th>
<th>Weighted Mean Difference (Fixed)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>Mean(SD)</td>
<td>N</td>
<td>Mean(SD)</td>
<td>95% CI</td>
</tr>
<tr>
<td>01 Strategy training</td>
<td>Donkervoort 2001</td>
<td>43</td>
<td>3.00 (4.10)</td>
<td>40</td>
<td>2.83 (3.20)</td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td>43</td>
<td>40</td>
<td>100.0</td>
<td>0.17 [ -1.41, 1.75 ]</td>
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<td></td>
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<td>Test for overall effect z=0.21 p=0.8</td>
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[Graph showing analysis results]

### Analysis 01.02. Comparison 01 Experimental therapy versus standard care, Outcome 02 Change in Barthel at end of therapy

Review: Interventions for motor apraxia following stroke  
Comparison: 01 Experimental therapy versus standard care  
Outcome: 02 Change in Barthel at end of therapy

<table>
<thead>
<tr>
<th>Study</th>
<th>Experimental</th>
<th>Control</th>
<th>Weighted Mean Difference (Fixed)</th>
<th>Weight (%)</th>
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<tr>
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<td>N</td>
<td>Mean(SD)</td>
<td>95% CI</td>
</tr>
<tr>
<td>01 Strategy training</td>
<td>Donkervoort 2001</td>
<td>45</td>
<td>2.44 (3.00)</td>
<td>48</td>
<td>1.15 (2.50)</td>
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<tr>
<td>Subtotal (95% CI)</td>
<td>45</td>
<td>48</td>
<td>93.9</td>
<td>1.29 [ 0.16, 2.42 ]</td>
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<td>Test for overall effect z=2.24 p=0.02</td>
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</table>

<table>
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<th>Experimental</th>
<th>Control</th>
<th>Weighted Mean Difference (Fixed)</th>
<th>Weight (%)</th>
<th>Weighted Mean Difference (Fixed)</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>Mean(SD)</td>
<td>N</td>
<td>Mean(SD)</td>
<td>95% CI</td>
</tr>
<tr>
<td>02 Transfer of training</td>
<td>Edmans 2000</td>
<td>3</td>
<td>4.00 (3.00)</td>
<td>6</td>
<td>2.80 (3.50)</td>
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<tr>
<td>Subtotal (95% CI)</td>
<td>3</td>
<td>6</td>
<td>6.1</td>
<td>1.20 [ -3.20, 5.60 ]</td>
<td></td>
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<td>Test for heterogeneity: not applicable</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Test for overall effect z=0.53 p=0.6</td>
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</table>

<table>
<thead>
<tr>
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<th>Weighted Mean Difference (Fixed)</th>
<th>Weight (%)</th>
<th>Weighted Mean Difference (Fixed)</th>
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<tbody>
<tr>
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<td>N</td>
<td>Mean(SD)</td>
<td>N</td>
<td>Mean(SD)</td>
<td>95% CI</td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td>48</td>
<td>54</td>
<td>100.0</td>
<td>1.28 [ 0.19, 2.38 ]</td>
<td></td>
</tr>
<tr>
<td>Test for heterogeneity: not applicable</td>
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<td>Test for overall effect z=2.31 p=0.02</td>
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<td></td>
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</tbody>
</table>

[Graph showing analysis results]
### Analysis 01.03. Comparison 01 Experimental therapy versus standard care, Outcome 03 Change in Barthel at 12 months after end of therapy

**Review:** Interventions for motor apraxia following stroke  
**Comparison:** 01 Experimental therapy versus standard care  
**Outcome:** 03 Change in Barthel at 12 months after end of therapy

<table>
<thead>
<tr>
<th>Study</th>
<th>Experimental</th>
<th>Control</th>
<th>Weighted Mean Difference (Fixed)</th>
<th>Weight</th>
<th>Weighted Mean Difference (Fixed)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N Mean(SD)</td>
<td>N Mean(SD)</td>
<td>95% CI (%)</td>
<td></td>
<td>95% CI (%)</td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td>0 0</td>
<td>0</td>
<td>0.0</td>
<td>Not estimable</td>
<td></td>
</tr>
</tbody>
</table>

- **Test for heterogeneity:** not applicable  
- **Test for overall effect:** not applicable

### Analysis 01.04. Comparison 01 Experimental therapy versus standard care, Outcome 04 Death

**Review:** Interventions for motor apraxia following stroke  
**Comparison:** 01 Experimental therapy versus standard care  
**Outcome:** 04 Death

<table>
<thead>
<tr>
<th>Study</th>
<th>Experimental</th>
<th>Control</th>
<th>Peto Odds Ratio</th>
<th>Weight</th>
<th>Peto Odds Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n/N</td>
<td>n/N</td>
<td>95% CI (%)</td>
<td></td>
<td>95% CI (%)</td>
</tr>
<tr>
<td>01 Strategy training</td>
<td>Dankervoord 2001</td>
<td>2/56</td>
<td>5/57</td>
<td>100.0</td>
<td>0.41 [ 0.09, 1.89 ]</td>
</tr>
<tr>
<td>Subtotal (95% CI) 56</td>
<td>57</td>
<td>100.0</td>
<td>0.41 [ 0.09, 1.89 ]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total events: 2 (Experimental), 5 (Control)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
| Test for heterogeneity: not applicable  
| Test for overall effect: z=1.14 p=0.3 |

| 02 Transfer of training | Edmans 2000 | 0/3 | 0/6 | 0.0 | Not estimable |
| Subtotal (95% CI) 3 | 6 | 0.0 | Not estimable |
| Total events: 0 (Experimental), 0 (Control) |
| Test for heterogeneity: not applicable  
| Test for overall effect: not applicable |

| 03 Gesture training | Smania 2000 | 0/6 | 0/4 | 0.0 | Not estimable |
| Subtotal (95% CI) 6 | 4 | 0.0 | Not estimable |
| Total events: 0 (Experimental), 0 (Control) |
| Test for heterogeneity: not applicable  
| Test for overall effect: not applicable |

Total (95% CI) 65 | 67 | 100.0 | 0.41 [ 0.09, 1.89 ] |

- **Test for heterogeneity:** not applicable  
- **Test for overall effect:** z=1.14 p=0.3

---

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Analysis 01.05. Comparison 01 Experimental therapy versus standard care, Outcome 05 Quality of life measures

Review: Interventions for motor apraxia following stroke
Comparison: 01 Experimental therapy versus standard care
Outcome: 05 Quality of life measures

<table>
<thead>
<tr>
<th>Study</th>
<th>Experimental</th>
<th>Control</th>
<th>Weighted Mean Difference (Fixed)</th>
<th>Weight</th>
<th>Weighted Mean Difference (Fixed)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean(SD)</td>
<td>N</td>
<td>95% CI (%)</td>
<td>(%)</td>
<td>95% CI (%)</td>
</tr>
<tr>
<td>Total (95% CI)</td>
<td>0</td>
<td>0</td>
<td>0.0</td>
<td>Not estimable</td>
<td></td>
</tr>
</tbody>
</table>

Test for heterogeneity: not applicable
Test for overall effect: not applicable
### Analysis 01.06. Comparison 01 Experimental therapy versus standard care, Outcome 06 Ability to gesture, pantomime, use real objects

Review: Interventions for motor apraxia following stroke  
Comparison: 01 Experimental therapy versus standard care  
Outcome: 06 Ability to gesture, pantomime, use real objects

<table>
<thead>
<tr>
<th>Study</th>
<th>Experimental</th>
<th>Control</th>
<th>Weighted Mean Difference (Fixed)</th>
<th>Weight</th>
<th>Weighted Mean Difference (Fixed)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N  Mean(SD)</td>
<td>N  Mean(SD)</td>
<td>95% CI (%)</td>
<td>95% CI</td>
<td></td>
</tr>
<tr>
<td>Gesture training</td>
<td>6  37.70 (16.20)</td>
<td>4  29.30 (20.90)</td>
<td>100.0</td>
<td>8.40 [-15.84, 32.64 ]</td>
<td></td>
</tr>
<tr>
<td>Subtotal (95% CI)</td>
<td>6</td>
<td>4</td>
<td>100.0</td>
<td>8.40 [-15.84, 32.64 ]</td>
<td></td>
</tr>
</tbody>
</table>
| Test for heterogeneity: not applicable  
Test for overall effect z=0.58  p=0.5

| Using real objects | 6  11.70 (2.30) | 4  10.50 (4.10) | 100.0 | 1.20 [-3.22, 5.62 ] |
| Subtotal (95% CI) | 6 | 4 | 100.0 | 1.20 [-3.22, 5.62 ] |
| Test for heterogeneity: not applicable  
Test for overall effect z=0.53  p=0.6

### Analysis 01.07. Comparison 01 Experimental therapy versus standard care, Outcome 07 Effects on family and carer

Review: Interventions for motor apraxia following stroke  
Comparison: 01 Experimental therapy versus standard care  
Outcome: 07 Effects on family and carer

<table>
<thead>
<tr>
<th>Study</th>
<th>Experimental</th>
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<th>Weight</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>N  Mean(SD)</td>
<td>N  Mean(SD)</td>
<td>95% CI (%)</td>
<td>95% CI</td>
<td></td>
</tr>
</tbody>
</table>
| Total (95% CI) | 0 | 0 | 0.0 | Not estimable

Test for heterogeneity: not applicable  
Test for overall effect: not applicable

---

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## Analysis 01.08. Comparison 01 Experimental therapy versus standard care, Outcome 08 Carer and family perceptions

**Review:** Interventions for motor apraxia following stroke  
**Comparison:** 01 Experimental therapy versus standard care  
**Outcome:** 08 Carer and family perceptions

<table>
<thead>
<tr>
<th>Study</th>
<th>Experimental</th>
<th>Control</th>
<th>Weighted Mean Difference (Fixed)</th>
<th>Weight</th>
<th>Weighted Mean Difference (Fixed)</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>Mean(SD)</td>
<td>N</td>
<td>Mean(SD)</td>
<td>95% CI</td>
<td>(%) 95% CI</td>
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<td>0.0</td>
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Test for heterogeneity: not applicable  
Test for overall effect: not applicable

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<tbody>
<tr>
<td>Favours control</td>
<td>Favours experimental</td>
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<td></td>
<td></td>
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</tbody>
</table>

## Analysis 01.09. Comparison 01 Experimental therapy versus standard care, Outcome 09 Economic resources

**Review:** Interventions for motor apraxia following stroke  
**Comparison:** 01 Experimental therapy versus standard care  
**Outcome:** 09 Economic resources

<table>
<thead>
<tr>
<th>Study</th>
<th>Experimental</th>
<th>Control</th>
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<th>Weight</th>
<th>Weighted Mean Difference (Fixed)</th>
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<tr>
<td>N</td>
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<td>Mean(SD)</td>
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<td>(%) 95% CI</td>
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Test for heterogeneity: not applicable  
Test for overall effect: not applicable

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<tbody>
<tr>
<td>Favours experimental</td>
<td>Favours control</td>
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</tbody>
</table>
### Analysis 01.10. Comparison 01 Experimental therapy versus standard care, Outcome 10 Apraxic patient's mood

**Review:** Interventions for motor apraxia following stroke  
**Comparison:** 01 Experimental therapy versus standard care  
**Outcome:** 10 Apraxic patient's mood

<table>
<thead>
<tr>
<th>Study</th>
<th>Experimental</th>
<th>Control</th>
<th>Weighted Mean Difference (Fixed)</th>
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<tr>
<td>N Mean(SD)</td>
<td>N Mean(SD)</td>
<td>95% CI</td>
<td>(%)</td>
<td>95% CI</td>
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</tr>
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</table>

- Total (95% CI): 0 0 0.0 Not estimable

Test for heterogeneity: not applicable  
Test for overall effect: not applicable

-10.0 -5.0 0 5.0 10.0  
Favours control Favours experimental