1	TITLE PAGE
2	Running Head: FES versus AFO for foot-drop
3	
4	Title: Functional electrical stimulation versus ankle foot orthoses for foot-drop: a meta-
5	analysis of orthotic effects
6	
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ABSTRACT

Objective: To compare the effects on walking of Functional Electrical Stimulation (FES) 45 and Ankle Foot Orthoses (AFO) for foot-drop of central neurological origin, assessed in 46 terms of unassisted walking behaviours compared with assisted walking following a 47 period of use (combined-orthotic effects). 48 Data Sources: MEDLINE, AMED, CINAHL, Cochrane Central Register of Controlled 49 Trials, Scopus, REHABDATA, PEDro, NIHR Centre for Reviews and Dissemination 50 51 and clinicaltrials.gov. plus reference list, journal, author and citation searches. 52 Study Selection: English language comparative Randomised Controlled Trials (RCTs). Data Synthesis: Seven RCTs were eligible for inclusion. Two of these reported different 53 54 results from the same trial and another two reported results from different follow up periods so were combined; resulting in five synthesised trials with 815 stroke 55 56 participants. Meta-analyses of data from the final assessment in each study and three 57 overlapping time-points showed comparable improvements in walking speed over ten metres (p=0.04-0.95), functional exercise capacity (p=0.10-0.31), timed up-and-go 58 59 (p=0.812 and p=0.539) and perceived mobility (p=0.80) for both interventions. *Conclusion:* Data suggest that, in contrast to assumptions that predict FES superiority, 60 AFOs have equally positive combined-orthotic effects as FES on key walking measures 61 for foot-drop caused by stroke. However, further long-term, high-quality RCTs are 62 required. These should focus on measuring the mechanisms-of-action; whether there is 63 64 translation of improvements in impairment to function, plus detailed reporting of the devices used across diagnoses. Only then can robust clinical recommendations be made. 65 Key words: electrical stimulation therapy, nervous system diseases, stroke, walking, foot 66 drop, systematic review, meta-analysis. 67

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MAIN TEXT

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INTRODUCTION

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73 Conditions such as stroke, brain injury (BI), multiple sclerosis (MS), spinal cord injury (SCI) and cerebral palsy (CP) affect upper motor neuronal pathways (1) and are collectively 74 referred to as pathologies of central neurological origin (CNO) (2). In the United Kingdom 75 (UK) there are approximately 1.2 million people living with stroke (3), 100,000 MS and 76 40,000 SCI (4), there are 160,000 BI admissions per year (5), and 1 in 400 people have CP 77 (6). Foot-drop is a common impairment seen across these conditions (7) and although 78 prevalence data in some of the CNO conditions is very limited, a commonly cited figure 79 80 suggests that it is seen in 20-30% of people with stroke (7, 8)81 Foot-drop is categorized as an inability to dorsiflex the foot, with or without excessive inversion and is most commonly caused by weakness in the dorsiflexor (and evertor) and/or 82 overactivity in the plantarflexor (and invertor) muscle groups. Foot-drop results in walking 83 being slower, less efficient and potentially unsafe (7); as foot clearance during swing and 84 initial foot contact at the start of the stance phase are compromised. These factors have been 85 associated with an increased risk of falls (7), reduced quality of life (7, 9) and increased 86

87 levels of mortality (10).

Current practice in the treatment of foot-drop normally involves a form of ankle foot orthosis
(AFO)(11). Functional electrical stimulation (FES) is also used but less frequently (9).

AFOs stabilise the foot and ankle and lift the toes when stepping (12). Meta-analyses have
shown them to have positive effects on some aspects of walking (12, 13) but these analyses
are primarily based on non-randomised control trial (RCT) evidence. AFOs have been
criticised for detrimental effects on the adaptability of walking, propulsion, aesthetics and
comfort (14-16) which can impact compliance and satisfaction.

Foot-drop FES uses electrical pulse trains to stimulate the common peroneal nerve over key
phases of the gait cycle to correct the foot-drop impairment (17). This phasic stimulation can
be delivered via surface or implanted electrodes. Foot-drop FES has been shown to have
positive effects on walking speed (18, 19) but meta-analyses have also, in part, been based on
non-RCT evidence. For surface systems, limitations have been cited in relation to issues with
effort of setup, skin irritation and pain (20), which again affects compliance and satisfaction.
Implanted systems address some of these limitations but are more costly (21).

Despite their limitations both are endorsed in the management of foot-drop with clinical 102 guidelines existing for AFO as a result of stroke (22, 23) MS (24), CP (25) and BI (26) and 103 104 FES guidelines promoting use across all CNO diagnoses (2). However, these guidelines have had to rely on some non-RCT sources of evidence and as intervention specific guidelines, 105 106 comparing to no treatment or physiotherapy, do not consider evidence from direct comparisons between these interventions. As a result current guidelines do not provide 107 108 clinicians with a clear patient pathway. Recently a number of RCTs providing direct comparisons have been published. Furthermore, these studies have advanced our 109 understanding of the effects these interventions may produce: 110

a) Immediate-orthotic effects where same-day comparisons are made between AFO/FES
unassisted and assisted walking behaviours (16, 27).

113	b) Therapeutic effects (19, 28) where unassisted walking behaviours are compared with
114	unassisted walking on a day some period later (16, 27).
115	c) Training effects (16) where assisted walking behaviours are compared with assisted
116	walking on a day some period later.
117	d) Combined-orthotic effects (15) where unassisted walking behaviours on one day are
118	compared with assisted walking on a day some period later (16, 27).
119	
120	The suggested mechanism-of-action for AFO is that the device remedies the loss of
121	dorsiflexion/eversion by holding the foot in a neutral position but this can result in negative
122	effects on neuromuscular control and muscle biomechanics with long-term use (29-31).
123	Therefore, it has been assumed that they only provide immediate-orthotic effects (a) (12), a
124	notion supported by the only known long-term AFO specific RCT in the field (32).
125	In contrast, there are many reports of long-term neuromuscular control improvements with
126	FES (19, 33) which are attributed to changes in neural plasticity, muscular strength and
127	cardiovascular efficiency (31, 34, 35). The mechanism for these improvements has been
128	hypothesised as being due to the coinciding of antidromic electrical stimulation-generated
129	action potentials with volitional activity leading to strengthening of modifiable Hebb-
130	synapses at a segmental level (34, 36, 37).
131	Given these proposed mechanisms-of-action it could be assumed that FES will provide a
132	distinct advantage over AFO with long-term use.
133	Two recent reviews (9, 38) have explored the long-term effects evidence for AFOs versus
134	FES in stroke survivors; both concluding that there was a preference for FES but insufficient
135	evidence to recommend one over the other. However, the first was not systematic (39) and
136	included non-RCT studies (9) and the other did not meta-analyse; possibly due to the breadth

137	of question posed (38). This review (38) reported that FES was superior at conserving energy
138	but included a paper where FES was combined with botulinum toxin (40) and another that
139	compared FES to therapy as opposed to AFO (41).
140	In order to provide improved clinical guidelines which will help clinicians determine which
141	of these interventions to prescribe and what the directly comparable effects are over a period
142	of use gold standard meta-analysis of RCT level evidence is required (42). Given that both
143	interventions are most commonly prescribed as long-term orthotics (9, 30) and the
144	assumption that studying long-term use will highlight any differences in walking behaviours
145	resulting from the different mechanisms-of-action we sought to perform a systematic
146	examination of the evidence base to address the question:
147	Are the combined-orthotic effects on walking for foot-drop of CNO greater for FES than
148	AFO?
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151	METHODS
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153	This review was designed according to the Preferred Reporting Items for Systematic Reviews
154	and Meta-analyses (PRISMA) statement (43). The full review protocol can be found at:
155	http://www.crd.york.ac.uk/PROSPERO/register_new_review.asp?RecordID=9892&UserID=
156	<u>6114</u>
157	Nine electronic databases were searched. These were MEDLINE (Ovid), AMED (Ovid),
158	CINAHL (EBSCO), Cochrane Central Register of Controlled Trials (CENTRAL), Scopus,
159	REHABDATA, PEDro, NIHR Centre for Reviews and Dissemination and clinicaltrials.gov.

A search strategy including controlled vocabularies related to "electric stimulation",
"walking" and "nervous system diseases" and terms such as "foot drop" and "electric*
stimulat*" were used with no date limits (full search strategy available on request from the
corresponding author). Reference list, citation, key author and journal searches were also
completed and all searches were limited to the English language.

165 Once duplicates were removed one reviewer (SP) screened titles and abstracts categorising each as 'possibly' or 'clearly not' relevant against the inclusion criteria (Table I). Full length 166 articles were retrieved for 'possibly relevant' studies and two unmasked reviewers (SP and 167 KH) independently assessed their eligibility (Table I) classing them as 'relevant', 'definitely 168 irrelevant' or 'unsure'. Different outcome measurements from the same trial reported in 169 separate publications were treated as a single publication; as were separate publications that 170 reported different data collection time-points within the same trial. Any disagreements or 171 'unsure' publications were discussed (between SP and KH). A third reviewer was available to 172 resolve any disagreements (LK). 173

174

175 Table I. Inclusion Criteria.

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SP extracted data using a predesigned proforma; trial details extracted related to the
characteristics of the included studies, participant and intervention details. Missing data
and/or aspects that required clarification were requested from trial authors (14, 16, 44, 45), by
SP (Appendix I). KH reviewed the extracted data for accuracy.

181 As an RCT-based review, and to avoid the limitations of scaled quality assessment tools (42,

182 46), the Cochrane risk of bias assessment tool (42) was used independently by two reviewers

183 (SP and KH) with a third reviewer (LK) available if necessary. To ensure impartiality, risk of

184 bias was based on published work only. Performance bias was not considered as the interventions precluded blinding of participants and measures were primarily objective (46). 185 Outcomes across the World Health Organisation's (WHO) International Classification of 186 Functioning, Disability and Health (ICF) (47) were extracted. This helped to identify if there 187 was any comparative evidence to support the assumed mechanisms-of-action and whether 188 189 they translated into function. Therefore, all measurements were categorised as either being within the body functions and structures (BFS), activity or participation domain (47) by SP, 190 using supporting literature (47-50). All post-intervention data collection point assisted-191 192 walking means and standard deviations (SD) were extracted with final-assessment data pooled for data analysis. Given the hypothesised mechanisms-of-action suggesting that FES 193 would have greater benefits than AFO with longer-term use; broadly overlapping time-point 194 195 data was also grouped for meta-analysis where possible. Standard errors were converted to SDs (14, 42, 51) and functional exercise capacity (an activity domain measurement (52)) was 196 considered as metres walked so was converted as necessary (15). 197

Meta-analyses were performed using RevMan 5.3® software. Where the same measurement 198 was used across more than two trials, outcomes were combined using mean difference (MD) 199 200 with 95% confidence intervals (CIs). Where an outcome was measured using different approaches, such as functional exercise capacity (distance walked in metres measured over 201 two, three or six minutes), standardised mean difference (SMD) with 95% CIs was used. For 202 crossover trials only pre-crossover data was extracted (15). Where there was more than one 203 arm looking at the same intervention the similarity at baseline to the other intervention and 204 size were used to decide which to use and the data from the most comparable group extracted 205 (15). 206

207	Heterogeneity was examined using visual inspection of forest plot, chi ² test and I ² statistic. If
208	the chi ² test showed heterogeneity which the I ² statistic identified as being moderate to low,
209	(<50% (42)) a fixed-effects model was used. A random-effects model was used for
210	heterogeneity of $>50\%$.
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213	RESULTS
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215	1836 citations were found of which seven were eligible for inclusion. Two of these reported
216	outcomes from the same participants (44, 53) so were grouped, and subsequently referred to
217	by the first publication date (44). One trial published results up to six months (14) and had
218	another publication reporting results at 12 months (51); so were also grouped. For meta-
219	analysis the relevant publication was used with the source identified by the date of the
220	publication on the corresponding forest plot. Thus a total of five RCTs, published between
221	2007 and 2015 with 815 participants, were available for meta-analysis (Fig. 1).
222	
223	<i>Fig. 1.</i> Flowchart of trial selection.
224	
225	Table II. Characteristics of included trials, participant and intervention details.
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228	Characteristics of included trials
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230	One trial used a multiple-site crossover design (15) with two AFO arms. Data from arm 2
231	(AFO-FES) was used as it was larger and similar to the FES group at baseline. The remaining
232	four trials used two arm parallel RCT design, two single-site (44, 45) and two multiple-site
233	(14, 16) (Table II).
234	
235	Participant details
236	
237	All the participants were over the age of 18 years and had suffered a stroke. Average time
238	since diagnosis ranged from 51.7 days (45) up to 6.9 years (14, 51). Of those trials that
239	reported hemiplegic side (16, 44, 45) there was a relatively even distribution (116:47.9%
240	right, 126: 52.1% left). Two of the trials recruited current AFO users (16, 44) whereas the
241	remaining four introduced the interventions to both groups for the first time (Table II).
242	
243	Intervention details
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245	Three of the trials (14-16, 51) reported providing "customized" AFOs prescribed by an
246	orthotist; plus a physiotherapist for Kluding et al (16). One used off-the-shelf AFOs (45)
247	which is appropriate practice with their, sub-acute, population (54) and one used a
248	combination (44). No trial reported any further details of the AFOs or how prescription
249	decisions were made; none were hinged. All-but-one study used surface FES systems (44),
250	one trial highlighted that "clinicians" setup FES for measurement (45) but no trial reported
251	details of setup parameters such as electrode placement, ramping, amplitude or frequency.
252	The setting where interventions were used varied with participants from three of the studies

253	using the devices within their own environment (14, 15, 44, 51). One trial used them in both
254	the participants own environment and under supervision (16) and one used them only under
255	supervision (45). All-day-use was encouraged in all-but-one of the trials (45), some with a
256	gradual introduction, although whether this was adhered to was not reported. Three trials
257	provided concurrent therapy for both groups (16, 44, 45) (Table II).
258	
259	Methodological Quality
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261	Table III Risk of Rias
201	Table III. Risk of Dius
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263	Table III summarises the quality assessment, Kluding et al (16) alone had no identified areas
264	of high risk of bias.
265	
266	Table IV Outcome measurements and intervention effects
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269	Outcome Measurements
270	All trials utilised ICF activity domain measurements; most commonly the 10-metre walk test
271	(Table IV). However, one did not collect any BFS domain measurements (14, 51) and
272	another lacked participation domain measurements (15). The intervention period studied
273	ranged from six weeks $(15) - 12$ months (51) .

274	To allow direct comparison of the assumed mechanisms-of-action and functional translation
275	the following results are presented according to ICF domains. The narrative comparison
276	found in Table IV is summarised below. Final-assessment meta-analyses are presented first.
277	There were three overlapping data time-points found at 4-6 weeks, 12-13 weeks and 26-30
278	weeks for activity domain measurements. These are categorised as short, medium and longer-
279	term respectively (Table IV); meta-analyses at these time-points are then presented.
280	
281	BFS
282	
283	Physiological cost index (PCI) (15), cadence (45), spatiotemporal/kinematics (44) and lower
284	limb Fugl-Meyer (16) were reported by single trials; therefore pooled-analysis was not
285	possible. All the trials found within-group improvements but no significant statistical
286	differences were reported for any of these measures by the primary authors except Kottink et
287	al (44) who found some spatiotemporal and kinematic differences in favour of FES (p <0.05)
288	(Table IV).
289	
290	Activity
291	
292	Final-assessment outcomes of 10-metre walking speed (all five trials, n=789) and functional
293	exercise capacity (three trials, n=761) were pooled. Meta-analysis showed between-group
294	comparable improvement (MD= 0.01, [-0.04, 0.05]; I ² =0%; p=0.79, <i>Fig. 2a</i>); and SMD -0.07
295	[0.22, 0.07], I ² =0%; p=0.31, <i>Fig. 3a</i>) respectively.

Fig. 2. Activity domain measurement: 10-metre (m) walk test metres per second (m/s)

299 *Fig. 3.* Activity domain measurement: functional exercise capacity metres (m).

300

The timed up-and-go test was used in two trials (16, 51), both reported between-group comparable improvement (p=0.812 and p=0.539), therefore meta-analysis was not required (Table IV).

All other final-assessment activity measures were used in single trials with between-groupcomparable improvement in all cases (Table IV).

306 Meta-analysis was possible for the 10-metre walk test using data at short (four trials, n=771),

medium (three trials, n=699) and longer-term (three trials, n=713) time-points (*Fig. 2b-d*). It

revealed comparable improvement in the short-term (MD= 0.02 [-0.05, 0.10]; I²=66%;

309 p=0.54, *Fig. 2b*)) and longer-term (MD= -0.00 [-0.04, 0.04]; I²=14%; p=0.95, *Fig. 2d*)). In

310 the medium-term there was a marginal, but significant, difference in favour of AFO (MD= -

311 0.04 [-0.09,-0.00]; I²=0%; p=0.04, *Fig. 2c*)).

Functional exercise capacity meta-analyses were performed for short (three trials, n=761) and

medium-term (two trials, n=692) time-points (Fig. 3b and c). Meta-analyses revealed

between-group comparable improvement (SMD= -0.12 [-0.26-0.02]; I²=0%; p=0.10, Figure

315 3b) and SMD= -0.10 [-0.25, 0.05]; $I^2=0\%$; p=0.19, *Fig.* 3c)).

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Participation

The mobility domain of the Stroke Impact Scale (SIS) was collected by three trials (n=701) (14, 16, 45). Meta-analysis showed between-group comparable improvement (MD 0.31 [-2.06, 2.68]; I^2 =41%; p=0.80, *Fig. 4*).

322

323 *Fig. 4.* Participation domain measurement: Stroke Impact Scale (mobility sub-scale).

324

325	Activity monitoring was used by two trials (16, 44) (Table IV) but their data collection
326	methods varied too significantly (steps taken compared to time spent in different positions) to
327	pool results. Kluding et al (16) found no significant differences in the number of steps taken
328	and Kottink et al (44) found the FES group spent significantly more time in sitting/lying than
329	the AFO group (p=0.04).
330	All other final-assessment participation measurements were used by a single trial (14) with
331	between-group comparable improvements found (Table IV).
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334	DISCUSSION
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336	This is the first systematic review, including meta-analysis, of studies comparing AFO to
337	FES as interventions for people with CNO foot-drop which focusses on the clinically relevant
338	combined-orthotic effects on walking. As a RCT-based review with meta-analysis guided by
339	the PRISMA statement (55) the results provide the highest level of evidence currently

340 available to support clinical decision making (42).

341	The RCTs were deemed to be of medium-methodological quality, which provides some
342	confidence in our results that both interventions demonstrate equal combined-orthotic
343	improvements in 10-metre walking speed, functional exercise capacity, timed-up-and-go and
344	the mobility sub-scale of the SIS; regardless of the length of time used.
345	Given the different hypothesized mechanisms-of-action detailed in the introduction it is
346	somewhat surprising that there was no differentiation between the two interventions for any
347	of the pooled measurements. To explore this result we examined outcome measurements
348	within the BFS domain (which directly reflect mechanisms-of-action (48)) and whether or not
349	these changes in BFS coincide with changes in activity and participation differentially
350	between the interventions and over different time-points of use.
351	
352	BES
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354	The majority of measurements used in the reviewed trials suggest that there are no
355	differences between the two interventions. However, given the suggestions of a negative
356	influence of AFO and a positive influence of FES on volitional muscle activation it was
357	surprising that none of the included trials reported electromyography (EMG) or strength data.
358	Throughout our systematic search of the literature we found only one RCT (which explored
359	therapeutic as opposed to combined-orthotic effects) which compared FMG activity between
360	therapeute as opposed to combined ortholic encets) which compared ENIG activity between
	FES and AFO treatments. This trial reported that EMG activity was greater following a
361	FES and AFO treatments. This trial reported that EMG activity was greater following a period of FES than AFO use (56).
361 362	FES and AFO treatments. This trial reported that EMG activity was greater following a period of FES than AFO use (56).Kottink et al (53) was the only reviewed trial to measure gait features and found differences
361 362 363	FES and AFO treatments. This trial reported that EMG activity was greater following a period of FES than AFO use (56).Kottink et al (53) was the only reviewed trial to measure gait features and found differences between a FES group and an AFO group. Despite these findings, that are supported by results

365 should capture such measurements to determine whether restorative as opposed to

366 compensatory changes are made (62) in order to more accurately understand the mechanisms-367 of-action.

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Activity & Participation

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Meta-analysis of three validated measures of the activity domain (49, 52) and one mobility 371 372 specific participation domain measurement (49, 52) indicate that AFOs and FES produce equivalent functional improvements to walking for people with foot-drop as a result of 373 stroke; regardless of length of use. The equivalency of effects between these interventions is 374 375 supported by non-RCT studies which have found no significant changes in activity domain measurements when FES is provided to AFO users (59, 60, 63). 376 377 Given the difference in hypothesized mechanisms-of-action between FES and AFO and the 378 lack of BFS measurements, the question remains as to how these comparable effects on activity/participation are achieved. One explanation is that both simply correct the 379 mechanical problem of foot-drop; as is suggested for AFO. However, this does not fully 380 explain the differences between immediate-orthotic effect and orthotic effect after a period of 381 use. The activity monitoring results from one trial highlight another potential explanation. 382 Kluding et al (16) found that the number of steps taken per day increased with use of either 383 intervention (1891-2069, AFO and 2092-2369, FES at six and 30 weeks). This increase in 384 385 repetition of walking in both FES and AFO intervention groups (facilitated by the correction of foot-drop) could explain the observed comparable improvements. Indeed intensity of task-386 specific repetition is widely accepted as critical for effective improvements of motor-387 388 impairments (64-66). This hypothesis is consistent with Kluding et al's suggestion that both

interventions achieve combined-orthotic effects through immediate-orthotic and trainingeffects (16).

A final hypothesis is that RCTs to date have not been long enough to detect differences given the predominantly chronic populations investigated (67). Bethoux et al (51) did not find differences at 12 months which may suggest even longer-term follow up is required (68). To facilitate comparisons all future trials should ensure that data collection time-points are justified against physiological processes underlying treatment effects.

This review had some limitations. Firstly, it has revealed that until 2007 research has been limited to examinations of a single intervention for a single diagnosis precluding comparisons between interventions which might usefully inform clinicians which intervention may be most suitable. Since 2007 comparative RCTs have been undertaken, making this review timely. Whilst future FES (9, 69) and AFO specific studies (13, 70, 71) are necessary for intervention development, where possible, research should be impairment focused in order to facilitate more discerning prescription.

403 Secondly, despite the literature search encompassing all CNO diagnoses, the reviewed trials only included participants who had experienced a stroke and who were over the age of 18 so 404 405 our results can only be applied to this population. Trials using different CNO populations are necessary given that current clinical guidelines encompass them. Similarly, in order to form 406 clinical guidelines indicating which subgroups of patients with any given CNO diagnosis 407 (e.g. time points post-stroke, severity of foot-drop impairment) might benefit most from 408 either intervention future studies with carefully defined inclusion/exclusion criteria are 409 needed. This approach is of critical importance in subsequent trials so that potentially 410 important clinical effects are not diluted in heterogeneous study groups. Until such a time as 411 sufficient high-quality RCTs in specific groups of patients become available any meta-412 analyses will also suffer similar limitations. 413

Thirdly, risk of bias was present in the reviewed studies with detection bias (assessor 414 blinding) the most common area. While this might impact our results this area of bias is 415 common within rehabilitation research. Indeed, previous FES (28) and AFO (12) reviews 416 have chosen to discount it, suggesting it is impractical to address in studies of medical 417 devices. It can also be argued that objective measures minimize the risk of this source of bias. 418 However, two trials (15, 16) attempted to control for this, suggesting that it is feasible to 419 420 blind assessors and should at least be considered in future trials (72). We based the quality assessment on published material alone; so as not to advantage trial authors who respond to 421 422 requests for additional data. Therefore a lack of reported methodological detail might account for some of the other unclear and high areas of bias found. 423 424 Finally, the reader should note that a range of different AFO and FES devices were used in 425 the included trials and our analysis combined these. While combining data from different 426 types of AFO/FES does not allow a detailed look at the possible different effects of each individual sub-type, assuming the prescription of devices within each trial was provided on 427 the basis of clinical judgement and complies with current guidelines, this allows for a 428 clinically relevant comparison. Furthermore, limited reports of the details of AFO and FES 429 interventions preclude reliable sub-group analyses. The traditional description of AFOs on 430 the basis of the material used (carbon fibre, plastic, metal) or mode of manufacture 431 432 (customized versus off-the-shelf (54) as with our included trials) should be discontinued. The 433 mechanical properties (stiffness, mass) of an AFO determine its behaviour (73) so it is these that should be measured and reported (73-75). Similarly, differences in outcome between 434 therapist and patient FES setup have been found (76, 77) so this should also be reported. 435 436 None of the included trials reported details of FES setup parameters and it remains unclear which set of parameters would be most useful when comparing across trials; further work is 437 438 required in this area.

439	In conclusion, despite very different hypothesised mechanisms-of-action for AFO and FES
440	this RCT, state-of-the-art review, with meta-analysis (39) conservatively indicates that AFOs
441	have positive combined-orthotic effects on walking that are equivalent to FES for foot-drop
442	caused by stroke. Methodological and reporting limitations within the current RCT pool
443	preclude clinical recommendations regarding which type of AFO or FES set-up to use for
444	particular patient groups from being made; as they do in guiding clinicians which
445	intervention to prescribe for a specific patient. However crucially, and for the first time,
446	barriers to achieving such clinical recommendations within research design and reporting
447	have been identified to progress future research. Furthermore long-term, high-quality RCTs
448	are required across CNO diagnoses. These should focus on measuring the mechanisms-of-
449	action, whether there is translation of improved impairment to function and reporting the
450	correct device details; only then will discerning prescription be possible.
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Design

• Randomised Controlled Trials (RCT)

Participants

• Participants with foot-drop of a central neurological origin

Intervention

- Common peroneal nerve FES to address the specific impairment of foot-drop, with or without other areas of stimulation
- Stimulation eliciting a muscular contraction
- Trials where common peroneal stimulation is used during walking (overground or treadmill) as part of the intervention
- Trials studying combined-orthotic effects of foot-drop FES
- Trials where foot-drop FES and another intervention are used in combination but foot-drop FES is measured independently

Comparator

• Trials comparing foot-drop FES with AFO (the term therapy was allowed as might involve AFO)

Outcomes

• Measures of walking

	Trial design	N Diagnosis (R):(L)	Men: Women	Age (years)	Time since diagnosis	Current or new AFO users	AFO	Mechanical properties reported	FES	Setup for measurement done by	Use
Bethoux (2014 & 2015)+	2 arm parallel Multiple sites	495 (242 FES: 253 AFO) CVA Not specified	FES=147:95 AFO=157:96	FES=63.87 (11.33) AFO=64.3 (12.01)	FES=6.9yrs (6.43) AFO=6.86yrs (6.64)	New	Customized	No	Surface Walkaide	Not specified	Home 2wk progressive wearing schedule then AD
Everaert (2013)*	3 arm crossover Multiple sites	78 (43 FES: 35 AFO) CVA Not specified	FES=32:6** AFO=19:12**	FES=57.1 (12.9)** AFO=55.6 (11.9)**	FES=6.4mos (3.8)** AFO=6.9mos (3.2)**	New	Customized	No	Surface Walkaide	Not specified	Home AD
Kluding (2013)+	2 arm parallel Multiple sites	197 (99 FES: 98 AFO) CVA 93:104	FES=51:48 AFO=67:31	FES=60.71 (12.24) AFO=61.58 (10.98)	FES=4.77yrs (5.29) AFO=4.34yrs (4.1)	Current	Customized*** PLUS TENS for 2wks	No	Surface NESS L300	Not specified	Both Bioness clinical protocols followed 15mins-AD Training: 15mins x2 day 1wk then 20mins 2xday next 2wks
Kottink (2007)*~	2 arm parallel Single site	29 (14 FES: 15 AFO) CVA 13:16	FES=10:04 AFO=10:05	FES=55.2 (11.36) AFO=52.87 (9.87)	FES=9.07yrs (9.29) AFO=5.67yrs (4.64)	Current	Combination***	No	Implanted 2-channel implant	Not specified	Home Gradual increase over 2wks, then AD
Salisbury (2013)†	2 arm parallel Single site	16 (9 FES: 7 AFO) CVA 10:6	FES=03:06 AFO=03:04	FES=55.8 (11.3) AFO=52.6 (17.2)	FES=51.7 days (34.6)	New	Off the shelf ***	No	Surface ODFS	Clinician for FES	Supervised Part of physiotherapy 20mins, 5 x wk with supervised/ independent walking as appropriate.

703 Table II. *Characteristics of included trials, participant and intervention details.*

Abbreviations: FES= functional electrical stimulation; AFO=ankle-foot orthosis; *=post intervention/dropout characteristics; +=ITT completed; ~=based on 2007 not 2012 data; †= Pre intervention/drop out

705 characteristics; CVA= Cerebrovascular accident/Stroke; ** post intervention/drop characteristics at later time point than is included in this review (12 weeks); yrs=years; mos=months; Customized= custom made/

706 modified AFO; Combination= Different AFOs used by different participants; off the shelf= prefabricated/unmodified AFO; ***= both groups continued with physical therapy alongside intervention; TENS=

707 transcutaneous electrical nerve stimulation with no motor response; wk=week; NESS L300=Bioness model; ODFS= Odstock foot-drop system; AD=all day.

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711 Table III. *Risk of Bias*.

		Random sequence generation (selection bias)	Allocation concealment (selection bias)	Blinding of outcome assessment (detection bias)	Incomplete outcome data (attrition bias)	Selective reporting (reporting bias)	Other bias
	Bethoux	U	Н	Н	L	L	L
	2014/2015						
	Everaert 2013	U	U	U	Н	L	L
	Kluding 2013	L	L	U	L	U	L
	Kottink 2007	H	U	Н	U	L	L
	Salisbury	Н	L	Н	U	L	L
712	2013 Abbreviations: L= Low	; U=Unclear; H=High.					
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724	Table IV. Outco	ome measurements an	d intervention effects.				

	Walking outcome measures used & ICF level	Outcome collection	Combined-orthotic effects
Bethoux et al (2014/2015+)	Activity: • 10MWT ¹ • 6min walk test (distance) • Gaitrite Functional Ambulation Profile+ • mEFAP (including TUG) Participation+: • SIS (Mobility, ADL/IADL & social participation domains combined) ¹ • SIS mobility sub-scale • Perry ambulation categories based on 10MWT results	0 Short:1mos (not published) Medium: 3mos (not published) Long:6mos 12 mos+	• FEST=AFOT
Everaert et al (2013)	BFS: PCI over 4min test ¹ Activity: 4min walking test (speed) ¹ 10MWT Modified RMI	0, 3wks Short: 6wks	 Modified RMI: between- group, post-intervention differences not reported FES¹=AFO¹: for other measures
Kluding et al (2013)	BFS: LL Fugl Meyer Activity: 10MWT (self and fast) ¹ TUG 6min walk test (distance) Participation: SIS mobility sub-scale Activity monitoring (Stepwatch ®)	0 Short: 6 weeks Medium: 12 weeks Long: 30wks (only change data published)	 FES[↑]=AFO[↑]
Kottink et al (2007)	BFS: • stride time* • stride length* • stride width* • step length* • stance phase %* • 1 st double support phase %* • 1 st single support phase %* • kinematics=hip, knee & ankle* Activity: • 10MWT • 6min walk (speed) • Speed* Participation: • Activity monitoring (ActivPAL®)	0 Long: 26wks	 FES>AFO: Longer 1st single support phase %*; shorter Stance phase; 1st double support phase %*; Speed*; 10MWT; 6min walk (speed) at 26 wks AFO spent less time less in sitting/lying than FES FES 1=AFO 1: all other measures

	Salisbury et al (2013)	BFS:		0	• $FES_{T} = AFO_{T}$
		•	Cadence (10MWT)	Short: 6wks	
		Activity:		Medium: 12wks	
		•	Speed (10MWT)		
		•	FAC		
		Participat	ion:		
		•	SIS mobility sub-scale		
726 727 728 729	Abbreviations: wks=week Activities of Daily Living (2012); FAC=Functional A than.	s; mos=mon Instrument Ambulation	ths; min(s)=minute(s); mEFAP=modified Emory Functional Ambulat al Activities of Daily Living; 10MWT=10-metre walk test; PCI=Physi categories; ¹ =identified as primary outcome measure by authors; += no	on Profile; TUG=Timed Up and Go ological Cost Index; RMI=Rivermea ot reported in Bethoux 2015 12 mont	QoL=Quality of Life; SIS=Stroke Impact Scale; ADL/IADL= d Mobility Index; BBS=Berg Balance Scale; *=from Kottink et al h follow up publication; 1=increase; >=greater than; = =equal to; <=less
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Fig. 1. Flowchart of trial selection.



		FES			AFO			Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
Bethoux 2015	0.647	0.312	242	0.659	0.318	253	54.2%	-0.01 [-0.07, 0.04]	
Everaert 2013	0.625	0.309	38	0.568	0.261	31	9.2%	0.06 [-0.08, 0.19]	
Kluding 2013	0.56	0.28	99	0.56	0.26	98	29.3%	0.00 [-0.08, 0.08]	
Kottink 2007	0.95	0.13	9	0.83	0.24	12	6.5%	0.12 [-0.04, 0.28]	
Salisbury 2013	0.35	0.15	3	0.5	0.45	4	0.7%	-0.15 [-0.62, 0.32]	
Total (95% CI)			391			398	100.0%	0.01 [-0.04, 0.05]	•
Heterogeneity: Chi ² =	3.34, df	= 4 (P =	0.50);	I ² = 0%					
Test for overall effect:	6 (P = 0.1	79)						Favours [AFO] Favours [FES]	

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760 2a) Final-assessment

		FES			AFO			Mean Difference		Mean Difference	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI		IV, Random, 95% CI	
Bethoux 2014	0.601	0.265	242	0.639	0.302	253	35.1%	-0.04 [-0.09, 0.01]			
Everaert 2013	0.625	0.309	38	0.568	0.261	31	18.2%	0.06 [-0.08, 0.19]			
Kluding 2013	0.53	0.25	99	0.54	0.25	98	30.7%	-0.01 [-0.08, 0.06]			
Salisbury 2013	0.31	0.1	5	0.12	0.14	5	16.0%	0.19 [0.04, 0.34]			
Total (95% CI)			384			387	100.0%	0.02 [-0.05, 0.10]		•	
Heterogeneity: Tau ^a = 0.00; Chi ^a = 8.91, df = 3 (P = 0.03); l ^a = 66% Test for overall effect: Z = 0.62 (P = 0.54)						= 66%			-0.5	-0.25 0 0.25 Favours [AFO] Favours [FES]	0.5

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2b) Short-term. Bethoux et al (2014) and Kluding et al (2013) data obtained via

763 correspondence with authors



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2c) Medium-term. Bethoux et al (2014) and Kluding et al (2013) data obtained via

766 correspondence with authors

		FES			AFO			Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
Bethoux 2015	0.647	0.312	242	0.659	0.318	253	60.2%	-0.01 [-0.07, 0.04]	
Everaert 2013	0.625	0.309	38	0.568	0.261	31	0.0%	0.06 [-0.08, 0.19]	
Kluding 2013	0.56	0.28	99	0.56	0.26	98	32.6%	0.00 [-0.08, 0.08]	+
Kottink 2007	0.95	0.13	9	0.83	0.24	12	7.2%	0.12 [-0.04, 0.28]	
Salisbury 2013	0.35	0.15	3	0.5	0.45	4	0.0%	-0.15 [-0.62, 0.32]	
Total (95% CI)			350			363	100.0%	0.00 [-0.04, 0.04]	. ◆
Heterogeneity: Chi ² =	2.33, df	= 2 (P =	0.31);	l ² = 149	6				
Test for overall effect: Z = 0.07 (P = 0.95)									-0.5 -0.25 U 0.25 0.5 Favours [AFO] Favours [FES]

- 768 2d) Longer-term. Kluding et al (2013) data from correspondence with authors
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Fig. 3. Activity measure: Functional exercise capacity metres (m).

		FES			AFO			Std. Mean Difference	Std. Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
Bethoux 2015	204.6	106.08	242	217.9	152.64	253	65.1%	-0.10 [-0.28, 0.08]	
Everaert 2013	109.44	49.2	38	102.72	42.96	31	9.0%	0.14 [-0.33, 0.62]	
Kluding 2013	189.25	114.99	99	197.64	96.42	98	25.9%	-0.08 [-0.36, 0.20]	
Total (95% CI)			379			382	100.0%	-0.07 [-0.22, 0.07]	•
Heterogeneity: Chi ² =	0.89, df=	2 (P = 0.	64); I ² :	= 0%					
Test for overall effect:	Z=1.01	(P = 0.31))						Favours [AFO] Favours [FES]

3a) Final-assessment. Kluding et al (2013) data obtained via correspondence with authors.



3b) Short-term. Bethoux et al (2014) and Kluding et al (2013) data obtained via

777 correspondence with authors

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			FES			AFO			Std. Mean Difference	Std. Mean Difference
	Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
	Bethoux 2014	207	115.44	242	216.3	111.3	253	71.6%	-0.08 [-0.26, 0.09]	
	Kluding 2013	181.38	100.56	99	195.78	95.09	98	28.4%	-0.15 [-0.43, 0.13]	
	Total (95% CI)			341			351	100.0%	-0.10 [-0.25, 0.05]	•
	Heterogeneity: Chi² =	0.15, df=	1 (P = 0.	70); l² =	:0%				-	
778	Test for overall effect	Z=1.32 ((P = 0.19))						Favours [AFO] Favours [FES]
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3c) Medium-term. Data obtained via correspondence with authors

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Fig. 4. Participation measure: Stroke Impact Scale (mobility sub-scale).

		FES			AFO			Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
Bethoux 2014	60.8	15.6	242	60.5	15.9	253	73.1%	0.30 [-2.48, 3.08]	
Kluding 2013	78.76	16.89	99	77.57	16.49	98	25.9%	1.19 [-3.47, 5.85]	
Salisbury 2013	65.08	10.74	7	86.11	15.71	2	1.0%	-21.03 [-44.21, 2.15]	
Total (95% CI)			348			353	100.0%	0.31 [-2.06, 2.68]	•
Heterogeneity: Chi ² =	3.39, df	= 2 (P =	0.18);	$l^2 = 419$	6			2	
Test for overall effect	Z = 0.25	(P = 0.	80)						Favours [AFO] Favours [FES]

791		APPENDIX I
792	Unpub	lished data
793	٠	Salisbury et al (45) published results were a combination of assisted and unassisted
794		walking data. On request assisted data was provided.
795	٠	Kluding et al (16) published change as opposed to post-intervention data, this was
796		provided on request.
797	•	Kottink et al (44) only displayed results from their 2007 study in graphical form and
798		did not respond to request for raw data.
799	•	Bethoux et al (14) published standard error, these were converted to SD (42).
800	•	Both Bethoux et al (14) and Kluding et al (16) provided unpublished time-point data
801		on request.
802	٠	Functional exercise capacity was converted from the speed (metres per second) for
803		Everaert et al (15).
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