

1 **Effects of intradialytic cycling exercise on exercise capacity, quality of life, physical**  
2 **function and cardiovascular measures in adult haemodialysis patients: a systematic**  
3 **review and meta-analysis.**

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24

25 **Abstract**

26 **Background.** Intradialytic cycling (IDC), delivered during haemodialysis (HD) has the  
27 potential to improve many health issues. This systematic review and meta-analysis examines  
28 the evidence on the effects of IDC on exercise capacity, quality of life (QOL), physical  
29 function & cardiovascular health.

30 **Methods.** Twenty-four databases were searched alongside internet, hand searching and  
31 consultation with experts. Eligibility criteria were cluster randomised, randomised and quasi-  
32 randomised controlled trials (RCTs) of IDC versus usual care in prevalent adult HD patients.  
33 Primary outcome measures were exercise capacity (VO<sub>2</sub> peak and field tests) and QoL.  
34 Secondary measures were cardiac and physical function.

35 **Results.** Thirteen RCTs were eligible. Eight provided data for use in meta-analyses, which  
36 indicated no significant change in VO<sub>2</sub> peak (MD 1.19 ml/kg/min , 95% confidence interval -  
37 1.15 to 3.52, p=0.3), physical (MC 1.97, 95% CI -8.27 to 12.22, p=0.7) or mental component  
38 (MC 3.37, 95% CI -7.94 to 14.68, p=0.6) summary scores of the Medical Outcomes Short  
39 Form 36, pulse wave velocity (MD -0.57m/s, -1.55 to 0.41, p=0.4), systolic (MD -  
40 2.28mmHg, -14.46 to 9.90, p=0.7) or diastolic blood pressure (MD 2.25mmHg, -3.01 to 7.50,  
41 p=0.4) following IDC. IDC, however, leads to an improvement in performance on the six-  
42 minute walk test (MD 87.84m 39.60 to 136.09, p=0.0004). All included studies were  
43 considered to have high risk of bias.

44 **Conclusions.** There is insufficient evidence demonstrating that cycling exercise during HD  
45 improves patient outcomes. High quality, adequately powered, RCTs of IDC are required.

46

47 **Keywords.** ESRD; haemodialysis; exercise; systematic reviews; meta-analysis.

48 **Introduction**

49 The incidence and prevalence of end-stage renal disease (ESRD) requiring dialysis is  
50 increasing, with the majority of patients undertaking haemodialysis (HD).(1) These patients  
51 have significantly increased morbidity and mortality, with cardiovascular disease the leading  
52 cause of death.(2) Skeletal muscle catabolism, malnutrition, anaemia, uraemia, chronic  
53 inflammation, co-morbidities, physical inactivity, together with ‘enforced’ sedentary time  
54 during HD(3) also contribute to a reduction in exercise and functional capacity which are  
55 associated with disability(4) increased healthcare utilisation(5) and reduced quality of life  
56 (QoL) (6).

57

58 Exercise interventions have the potential to target several of these issues. The majority of  
59 previous reviews have examined the effects of exercise in general within the HD population  
60 (7-10). ‘Intradialytic exercise’ (an umbrella term covering a range of heterogeneous exercise  
61 interventions delivered during HD) is, however, often advocated due to greater adherence  
62 rates (11). Recent systematic reviews indicate that intradialytic exercise can significantly  
63 improve exercise capacity (12), physical QoL(12, 13) and blood pressure(13), but the  
64 interventions used include a range of different components. In choosing to review methods of  
65 exercise delivery rather than the specific type of intervention undertaken, clinicians and  
66 policy makers lack clear information on which specific modes of IDE are most beneficial,  
67 hampering the translation of research evidence into practice.

68

69 As intradialytic exercise delivered solely by means of a cycle ergometer (intradialytic  
70 cycling, IDC) is most commonly delivered within clinical practice(14) the aim of this review  
71 is to provide an up-to date synthesis of available evidence comparing the effects of IDC

72 versus usual care on exercise capacity, QoL, physical function, and cardiovascular health in  
73 HD patients.

74

## 75 **Materials and methods**

### 76 *Protocol registration and eligibility criteria*

77 A pre-specified protocol was published on PROSPERO ([www.crd.york.ac.uk/PROSPERO](http://www.crd.york.ac.uk/PROSPERO)  
78 [identifier CRD42016030006](https://doi.org/10.1111/1365-2702.12345)). Eligibility criteria included cluster randomised, randomised  
79 and quasi-randomised controlled trials (RCT) of prevalent HD patients. Trials in which it was  
80 possible to elucidate the direct effects of IDC training were included; studies of acute  
81 responses to a single bout of IDC were excluded. Control groups were those who received  
82 usual care haemodialysis, defined as four hours thrice weekly, not receiving any form of  
83 exercise intervention, counselling or education. Primary outcomes measures were: exercise  
84 capacity measured by VO<sub>2</sub> peak; treadmill or field tests (e.g. 6MWT or Incremental Shuttle  
85 Walk Test (ISWT) and quality of life (QoL) using validated measures. Secondary outcome  
86 measures included: cardiac (echocardiogram, blood pressure and pulse wave velocity) and  
87 physical function determined using a range of measures (Sit to Stand 5, STS5; Sit to Stand  
88 30, STS30; Sit to Stand 60, STS60; North Staffordshire Royal Infirmary Walk Test, NSRI;  
89 Timed Up and Go Test, TUAG).

90

### 91 *Study Identification*

92 To identify existing relevant trials and systematic reviews, the pre-specified databases, in  
93 addition to the National Institute of Health Research (NIHR) Centre for Reviews and  
94 Dissemination Database, SciELO (Scientific Electronic Library Online), OAIster (Open

95 Archives Initiative), SCOPUS, BASE (Bielefeld Academic Search Engine) and Open Grey,  
96 were searched from inception to March 2017, supplemented with internet searching until July  
97 4th, 2017, hand searching reference lists and consultation with experts. No restrictions were  
98 placed upon publication, language status, or year of publication. Search terms were adapted  
99 to database requirements, the full search strategy for the MEDLINE database is shown in  
100 supplemental item S1.

101

102 *Study selection, data extraction and risk of bias assessment*

103 Search results were managed using Refworks (ProQuest, Ann Arbor, MI, USA). Duplicate  
104 citations were removed, and the remaining citations screened against eligibility criteria. Titles  
105 and abstracts deemed not to meet these criteria were excluded. For the remaining citations,  
106 full-text articles were assessed for eligibility by HMLY, DSM and MG. Disagreements were  
107 resolved by discussion with recourse to a fourth reviewer if needed.

108

109 Data from eligible studies were extracted by one reviewer and checked by another, using a  
110 template based upon the Cochrane collaboration tool. Information relating to study design,  
111 population, intervention, usual care control, outcomes and adverse events were recorded.  
112 Where multiple reports originated from a single study, comparison of the key characteristics  
113 were made using tables and were only included, as a single study, if they reported relevant  
114 outcomes to avoid overstating results. Where missing data was encountered, original authors  
115 were contacted.

116

117 Reviewers independently assessed study quality according to the Cochrane Risk of Bias  
118 assessment tool. Risk of bias for randomized controlled trials was assessed as high, low, or  
119 unclear across five domains (15). The overall risk of bias was determined using the following  
120 criteria: 1. Low risk of bias (all criteria graded adequate), 2. Moderate risk of bias (one  
121 criterion graded inadequate or two unclear) and 3. High risk of bias (more than one criterion  
122 graded inadequate or more than two unclear).

123

#### 124 **Statistical analysis**

125 Descriptive statistics were used for characteristics of included studies. Where means and SD  
126 were neither reported nor available from study authors (16, 17), they were estimated from  
127 medians and IQRs(18) or standard deviations were calculated from standard error (19, 20).  
128 Only data from the first period within cross-over trials was used to reduce the potential  
129 influence of carry-over.

130

131 All outcomes were treated as continuous data and interpreted as mean differences. Analysis  
132 was primarily based upon final values. In circumstances of baseline imbalances between  
133 groups, analysis was based upon changes from baseline(15). Statistical heterogeneity was  
134 assessed with the  $I^2$  test, which describes the percentage of variation across studies above that  
135 attributed to chance(21). Heterogeneity was considered unimportant for  $I^2$  values up to  
136 40%(15). RevMan 5.3 software was used to undertake meta-analyses. Data pertaining to  
137 similar outcome measures were pooled (with participant as the unit of analysis) in a meta-  
138 analysis, using a random effects model, which assumes the observed estimates of treatment  
139 effect vary across studies due to within and between study variance (22, 23). Pre-planned

140 subgroup and sensitivity were not possible due to the limited number of studies available for  
141 meta-analysis and all studies being classified as high risk of bias respectively.

142

## 143 **Results**

### 144 *Search results and study characteristics*

145 Figure 1 provides a flow diagram of included studies. Searching identified 3669 following  
146 removal of duplicates. Following screening of titles, abstracts and full text articles. 13 studies  
147 were eligible for inclusion. A table detailing the excluded studies can be found in  
148 supplemental item S2. Due to inadequate reporting of group sizes, results and wide  
149 heterogeneity of measures only eight trials provided information for use in meta-analyses (17,  
150 19, 20, 24-28).

151 Supplementary item S3 provides a summary the characteristics of included trials. Included  
152 trials were published between 1995 and 2016 in English. Twelve were individually  
153 randomised(17, 19, 20, 24-33), and one cluster randomised (16). Twelve used a parallel  
154 group design and one a crossover design (17). All trials measured outcomes at baseline and at  
155 the end of the intervention(16, 17, 19, 20, 24-33), two also measured outcomes at an interim  
156 time-point(27, 33). Only one study included longer follow up, one month post intervention  
157 (28). In total, 369 patients were randomised to receive IDC (n=190) or usual care (n=179)  
158 with sample sizes ranging from 18-49.

159

### 160 *Characteristics of IDC*

161 Full reporting of the IDC intervention was lacking in several studies, and the characteristics  
162 of IDC interventions ranged widely between studies. Five studies did not provide information

163 regarding the mode of IDC training used(17, 24, 26, 29, 30, 32), four provided continuous  
164 training(19, 20, 27, 31), two interval training(16, 33) and two a combination(25, 28). Mean  
165 duration of exercise training was 14 weeks (range 6-26). In all but one study, where patients  
166 exercised twice a week (16), patients were encouraged to exercise at each HD session. The  
167 mean duration of each planned bout of training was 31 minutes (range 20-45) and the  
168 majority required patients to exercise at a moderate intensity(19, 20, 24-27, 29, 33). Actual  
169 levels of concordance were difficult to ascertain, as only four studies reported adherence level  
170 (17, 25, 26, 28, 29) and only three summarised the amount of exercise achieved (17, 26, 29,  
171 33). Where reported, mean adherence rates were 81%. Only five trials reported adverse  
172 events (16, 17, 25-27, 29) of which four reported no events (16, 17, 25, 27), and one no  
173 ‘significant complications’(26, 29). Exercise training was provided by researchers in three  
174 studies(20, 25, 27), “physical activity experts” in one study (19) and HD staff in another (17).  
175 One study reported the intervention being provided by medical and nursing staff, but it was  
176 unclear whether these were attached to the trial or the HD unit (31). Seven did not report who  
177 was delivering the intervention (16, 24, 26, 28-30, 32, 33).

178

### 179 **Risk of bias assessment**

180 Risk of bias summaries for all included studies are provided in table 1. All studies were rated  
181 as high risk of bias, primarily due to insufficient reporting. Only one study provided the  
182 information required to assess bias across all domains and was judged to be of high risk in all  
183 (25).

184

### 185 *Effect of IDC*

#### 186 *Exercise capacity*

187 Nine studies reported outcomes related to exercise capacity. Six studies measured VO<sub>2</sub>  
188 peak(16, 19, 26-28), however it was not possible to include VO<sub>2</sub> data from the cluster RCT in  
189 meta-analyses due to inadequate reporting (16). One study used maximum work capacity,  
190 measured in watts (33). Three studies used the 6MWT (19, 25, 31) and one the ISWT (20),  
191 both field tests of exercise capacity. Four studies including 85 participants provided VO<sub>2</sub> peak  
192 data appropriate for meta-analysis. A non-significant improvement of 1.19 ml/kg/min (95%  
193 confidence interval -1.15 to 3.52; p=0.3) figure 2A) was observed immediately following  
194 IDC compared to usual care (19, 26-28). Statistical heterogeneity was low I<sup>2</sup>=10%. The study  
195 that reported maximal work capacity also reported no significant difference between IDC and  
196 usual care (33).

197

198 Two of the three studies that assessed 6MWT provided sufficient information to pool results  
199 (19, 25). These trials included 48 participants and demonstrated a statistically significant  
200 improvement of 87.84m (39.60 to 136.09, p=0.0004, figure 2B) in favour of IDC. There was  
201 no evidence of statistical heterogeneity (I<sup>2</sup>=0%). The study excluded from synthesis also  
202 suggested a significant improvement in the IDC group (p<0.05) (31). Only one included  
203 study assessed ISWT (20) whereby a statistically significant 15% (p=0.03) improvement in  
204 distance walked was observed in the IDC group.

205

### 206 *Quality of life (QoL)*

207 The SF-36 tool reports QoL as physical (PCS) and mental component scores (MCS).(34)  
208 Two studies with a total of 52 participants reported the effect of IDC on PCS and MCS. Due  
209 to baseline imbalances between groups for one of the two included studies (25) both mental  
210 and physical component summary scores for the SF-36 were assessed as change from

211 baseline scores. Of these studies, only one reported mean change and standard deviation.(24)  
212 For the remaining study (25) we used a conservative estimated correlation coefficient value  
213 of 0.5(35) to calculate SD of change scores. No statistical differences were observed between  
214 IDC and usual care for the PCS (1.97, -8.27 to 12.22, p=0.7, figure 3A) or MCS scores (3.37,  
215 -7.94 to 14.68, p=0.6, figure 3B) of the SF-36. Statistical heterogeneity was not evident for  
216 the MCS ( $I^2=0\%$ ), and was negligible for the PCS ( $I^2= 18\%$ ).

217

### 218 *Cardiac outcomes*

219 Five studies reported cardiovascular outcomes, of these three measured resting systolic (SBP)  
220 and diastolic blood pressure (DBP)(17, 20, 25), two measured pulse wave velocity(17, 25).  
221 Three studies recorded findings from echocardiogram (20, 29, 30).

222

223 One trial was excluded from meta-analysis for systolic blood pressure (SBP) due to a large  
224 baseline difference of 21.4mmHg between the usual care and IDC groups (20). This study did  
225 not report any significant effect on BP results. Synthesised data from the remaining two trials  
226 (17, 25), including 50 participants revealed a non-significant reduction in SBP of -  
227 2.28mmHg with IDC (-14.46 to 9.90, p=0.7, figure 4A). Synthesised data from three trials  
228 including 67 participants demonstrated a non-significant increase of 2.25mmHg in diastolic  
229 blood pressure with IDC (-3.01 to 7.50, p=0.4, figure 4B). There was no evidence of  
230 statistical heterogeneity for the SBP ( $I^2=0\%$ ) and negligible heterogeneity for the DBP  
231 ( $I^2=2\%$ ).

232

233 Synthesised data from two trials measuring pulse wave velocity (17, 17, 25) including 50  
234 participants demonstrated a non-significant improvement of -0.57 m/s (-1.55 to 0.41, p=0.4,

235 figure 4C in the IDC group. There was no evidence of statistical heterogeneity ( $I^2=0\%$ ). Of  
236 the three trials reporting echocardiography measures, two trials, including 37 participants  
237 provided results for left ventricular mass index (LVMI). However, we deemed meta-analysis  
238 inappropriate due to a large, clinically relevant baseline difference between the control and  
239 IDC groups of  $28.3\text{g/m}^2$  in one study (20) and  $13.9\text{g/m}^2$  in the other (29). Neither reported  
240 any significant difference in LVMI following IDC.

241

242 Two studies including 62 patients measured left ventricular ejection fraction (LVEF). These  
243 were not included in a meta-analysis due to inadequate reporting of group size in one study  
244 (30). This study saw a significant improvement in LVEF in the IDC group ( $p=0.004$ )(30)  
245 whilst the other reported no significant change (29).

246

#### 247 *Physical function*

248 Three studies reported physical function outcomes. The clinical diversity in outcome  
249 measures prevented pooling of results, but inspection of individual studies revealed limited  
250 impact of IDC across a range of measures. A single study of 29 participants revealed no  
251 statistically significant change in TUAG scores (-0.3 seconds, -1.39 to 0.77,  $p=0.9$ ) between  
252 IDC and usual care (25). Similarly, a single trial of 22 participants that reported a number of  
253 function measures observed non-statistically significant improvements in STS5 (-0.57  
254 seconds, -1.85 to 0.71,  $p=0.4$ ), STS30 (0.59 stands, -2.76 to 3.94,  $p=0.7$ ), STS60 (0.50 stands,  
255 -5.67 to 6.67,  $p=0.9$ ), normal gait speed (0.43 seconds, -0.67 to 1.53,  $p=0.5$ ) and fast gait  
256 speed (0.08 seconds, -0.41 to 0.57,  $p=0.8$ ) (24). A baseline difference of 31.98 seconds  
257 between the IDC and control group was noted for the NSRI walk test, but analysis of mean  
258 change scores revealed a statistically significant improvement in the NSRI walk test (-10.2

259 seconds, -17.6 to -2.8,  $p=0.007$ ) following IDC (24). Finally, a single trial of 21 participants  
260 reported a significant improvement in the number of steps completed in six minutes in the  
261 IDC group compared with usual care (32), however a clinically relevant baseline imbalance  
262 of 15 steps between groups may have influenced these results.

263

## 264 **Discussion**

### 265 **Main Findings**

266 The results of this systematic review and meta-analysis suggest that IDC can lead to  
267 statistically and clinically significant improvements in exercise capacity measured via field  
268 testing, but current evidence demonstrates no statistically significant effect upon  $VO_2$  peak,  
269 QoL, blood pressure or arterial stiffness. Due to issues with trial reporting, substantial  
270 baseline imbalances in outcome measures between intervention groups and inconsistency in  
271 selection of outcome measures across studies, it was not possible to synthesise the effect of  
272 IDC on cardiac or physical function. The quality of the included evidence was considered to  
273 be at high risk of bias. Based upon this review, there is currently insufficient evidence to  
274 support the use of IDC in clinical practice to improve exercise capacity, QoL, cardiac or  
275 physical function.

276

### 277 **Comparison to previous reviews**

278 Our review, which focussed specifically upon intradialytic aerobic exercise, delivered using a  
279 static cycle ergometer, stands predominantly in contrast to previous publications. These  
280 reviews have shown significant improvements in exercise capacity (12, 13) physical QoL(12)  
281 and blood pressure (13). These differences may be explained by the inclusion of additional  
282 evidence in our data synthesis, but also by the decision of previous reviews to include

283 intradialytic exercise programmes consisting of a range of types of exercise within their  
284 meta-analyses and interventions that were not exclusively aerobic training(36, 37) or  
285 delivered to HD patients (38) within subgroup analyses investigating types of intradialytic  
286 exercise modality (13).

287

## 288 **Implications for practice**

289 Current evidence suggests patients, on average, observe a clinically important improvement  
290 of 87m in the 6MWT after a programme of IDC, although the strength of this conclusion is  
291 limited by the high risk of bias within the two included studies (39). Theoretically an  
292 improvement in field tests of exercise capacity might reasonably be expected to be reflected  
293 in improvement in VO<sub>2</sub> peak, given that aerobic capacity contributes towards overall exercise  
294 capacity. The contrasting results seen in these two outcomes may be due to the inclusion of  
295 greater numbers of different trials within the meta-analysis for VO<sub>2</sub> peak in comparison to  
296 that for the 6MWT, other factors (e.g. muscular fatigue, anaerobic threshold, motivation)  
297 known to influence field test performance(5)andthe use of different protocols to measure  
298 exercise capacity, which can influence the measurement of VO<sub>2</sub> peak, and its sensitivity to  
299 change following a programme of exercise(40) The high prevalence of autonomic  
300 dysfunction in many HD patients may also be another reason for the discrepancy between the  
301 results of the two outcomes (41). Autonomic dysfunction can lead to cardiac  
302 unresponsiveness which is in turn associated with poor physical performance. Whilst  
303 speculative, it is possible that this may play a more important role during a higher intensity  
304 exercise test and therefore, patients may achieve greater improvements in field tests such as  
305 the 6MWT, when compared with VO<sub>2</sub> peak. Although, reduced exercise capacity and  
306 sedentary behaviour are powerful predictors of mortality in end-stage renal disease (4, 42,

307 43), these results should not be extrapolated, as a causal link between increased exercise  
308 capacity through exercise training and decreased mortality has yet to be established. IDC  
309 appeared also to have little influence upon cardiovascular outcomes within the current  
310 review, although this may reflect a lack of high quality RCTs rather than evidence of  
311 ineffectiveness.

312

313 Previous reviews have reported no significant adverse events (9, 12, 13) due to exercise  
314 training, citing these as evidence of safety. In the current review, eight trials failed to report  
315 information on adverse events and those which did only provided limited information.

316 Therefore, it is difficult to make a clear judgement about the safety of IDC. A recent study of  
317 IDC noted a significant drop in blood pressure one hour post exercise, with no reported  
318 adverse events (44). Given the association between asymptomatic intradialytic hypotension  
319 and adverse outcomes (45, 46), further research is needed to confirm that IDC is not  
320 associated with subclinical adverse events or to provide clinicians with information about  
321 groups of patients for whom IDE is not appropriate.

322

323 Exercise is beneficial across a spectrum of chronic diseases(47) and current Kidney Disease  
324 Outcomes Quality Initiative guidance recommends patients with chronic kidney disease CKD  
325 undertake “an exercise program compatible with cardiovascular health and tolerance, aiming  
326 for at least 30 minutes 5 times per week” (48). Expert statements provide more specific  
327 guidance (49, 50) and systematic reviews have extolled the benefits of exercise for HD  
328 patients (7-10, 12, 13). These have all led to increasing calls for exercise, particularly  
329 intradialytic, to become a routine part of the care of HD patients (13, 51, 52). The results of  
330 this review suggest, however, that the specific effects of IDC are currently unknown,

331 primarily due to the methodological shortcomings of existing trials. Further high quality,  
332 well-reported, RCTs are required. Two such trials, PEDAL (NCT02222402) and CYCLE-HD  
333 (ISRCTN11299707), will imminently provide evidence of the ability of intradialytic exercise  
334 to influence cardiac function and quality of life, which may further inform best practice.

### 335 **Implications for research**

336 Table 2 outlines several implications for future research which have been highlighted by this  
337 review. Unclear reporting was a feature of most IDC trials, and the primary reason for high  
338 risk of bias. Evidence presented by unclear trials rarely leads to change in practice or  
339 advances in research, because clinicians and policy makers lack confidence in the validity of  
340 findings and interventions can rarely be replicated (53, 54). Whilst we acknowledge that  
341 several IDC trials included within this review precede the publication of this guidance, future  
342 trials should adhere to both CONSORT guidance for the reporting of non-pharmacologic  
343 trials and use the Template for Intervention Description and Replication (TIDieR) checklist  
344 for intervention reporting (54, 55).

345

346 Most trials were small, and not powered to detect a true effect (56). Studies reporting  
347 echocardiography and harms are particularly vulnerable to lack of statistical power (57-60).  
348 Lack of blinding, which can influence participant, intervention provider and outcome  
349 assessor behaviour, leading to over-estimation of effect(56, 61, 62) particularly within non-  
350 pharmacologic trials (63) and ‘per protocol’ analysis, which may bias the estimated effects,  
351 were also common (61, 64, 65). Future trials should endeavour to adequately power studies,  
352 and aim to report blinding methods and fidelity explicitly (56).

353

354 Inconsistent use of a wide range of outcome measures limited meta-analyses, highlighting the  
355 need for a core outcome set to be measured and reported in all trials, alongside outcomes  
356 relevant to the individual study. Standardised Outcomes in Nephrology (SONG) aims to  
357 develop a core set of validated outcomes that reflect the main concerns of key stakeholders,  
358 including patients (66). Thirty-four priorities areas have been identified, approximately 21 of  
359 which may potentially be influenced by exercise interventions. Once standardised outcome  
360 measures have been established, better synthesis of studies will be possible, allowing  
361 comparisons of different interventions across outcomes (66, 67).

362

### 363 **Limitations**

364 Due to inadequate reporting two potentially relevant studies were excluded (38, 68).  
365 Additionally, six included trials presented outcome data incompletely (16, 20, 27, 31-33).  
366 Attempts to obtain these data were unsuccessful. The inclusion of these may have provided  
367 additional information for meta-analyses, potentially providing larger sample sizes and  
368 greater statistical power. The limited number of eligible studies meant it was not possible to  
369 assess publication bias or conduct subgroup analyses. Further analyses of specific  
370 characteristics of the IDC intervention (e.g duration of programme, adherence, intensity of  
371 exercise) would have provided information on potential reasons for differences in outcomes.  
372 Despite the lack of funnel plots, risk of publication bias is minimal, as many included studies  
373 reported statistically non-significant results. Unpublished studies with statistically non-  
374 significant results may exist, but their addition is not likely to change our conclusions.

375

### 376 **Conclusions**

377 The renal community remains in a position of equipoise regarding IDC, because of a lack of  
378 high quality RCT data. This review highlights the need for adequately powered trials that  
379 adhere to published reporting guidance and, as far as possible, take steps to remedy the  
380 methodological limitations of trials that have gone before.

381

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387

### 388 **Conflicts of Interest Statement**

389 All authors declare that they have no conflicts of interest. The results presented in this paper  
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391

392 **Contributions:** Conception; HMLY, SJS, ACS, JOB. Designed the protocol; HMLY.  
393 Assisted with protocol design; DSM, MGB, AJ, FC, SJS, CB. Co-ordinated the review  
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<b>Study</b>	<b>Random sequence generation</b>	<b>Allocation concealment</b>	<b>Blinding of participants</b>	<b>Blinding of personnel</b>	<b>Blinding of outcome assessment</b>	<b>Incomplete outcome data</b>	<b>Reporting bias</b>	<b>Other bias</b>	<b>Overall risk</b>
<b>Carmack 1995</b> (28)	Unclear	Unclear	Unclear	Unclear	Unclear	Unclear	Unclear	Low risk	High risk
<b>DeLima 2013</b> (32)	High risk	Low risk	Unclear	Unclear	Unclear	Low risk	High risk	High risk	High risk
<b>Giannaki 2013</b> (16, 16, 24)	Unclear	Unclear	Unclear	Unclear	High risk	Low risk	Unclear	Low risk	High risk
<b>Groussard 2015</b> (19)	Unclear	Unclear	High risk	High risk	High risk	Low risk	Unclear	Low risk	High risk
<b>Koh 2010</b> (25)	Low risk	Low risk	High risk	High risk	High risk	High risk	High risk	High risk	High Risk
<b>Liao 2016</b> (31)	Unclear	Unclear	High risk	High risk	High risk	Low risk	High risk	Low risk	High risk
<b>Momeni 2014</b> (30)	Unclear	Unclear	Unclear	Unclear	Unclear	Unclear	Unclear	Low risk	High risk
<b>Moug 2004</b> (16)	Unclear	High risk	Unclear	Unclear	Unclear	Unclear	Unclear	Low risk	High risk
<b>Painter 2002</b> (27)	Low risk	Unclear	Unclear	Unclear	High risk	Unclear	High risk	Low risk	High risk
<b>Parsons 2004</b> (33)	Low risk	Unclear	Unclear	Unclear	Unclear	High risk	Unclear	Low risk	High risk
<b>Reboredo 2010 and 2011</b> (26, 29, 29)	Unclear	Unclear	Unclear	Unclear	Unclear	High risk	Unclear	High risk	High risk
<b>Toussaint 2008</b> (17)	Unclear	Low risk	Unclear	Unclear	Unclear	Low risk	Unclear	Low risk	High risk
<b>Wilund 2010</b> (20)	Unclear	Unclear	Unclear	Unclear	Unclear	Low risk	High risk	High risk	High risk

611 **Table 1. Risk of bias for included studies.**

<b>Aim</b>	<b>Methodological limitation</b>	<b>Rationale for inclusion</b>	<b>Potential strategies to address limitation</b>
<b>Improved reporting of trials</b>	Poor reporting of delivery of intervention	Seven studies did not report intervention providers. Four studies reported adherence levels and three summarised amount of exercise data. Five studies did not provide any information regarding the type of IDC training used	<ul style="list-style-type: none"> <li>- Adhere to CONSORT guidance for the reporting of non-pharmacologic trials(54, 55);</li> <li>- Use Template for Intervention Description and Replication (TIDieR) checklist for intervention reporting.</li> </ul>
	Poor reporting of study design and trial procedures	All studies were rated as high risk of bias, primarily due to lack of sufficient reporting.  Nine trials provided insufficient details and allocation concealment.	<ul style="list-style-type: none"> <li>- Adhere to CONSORT guidance for the reporting of NPT.(53)</li> </ul>
	Poor reporting of testing procedures in relation to field testing and patient reported outcomes	Two studies provided no information on the procedures used for testing of field tests.  Of those reporting, use of a familiarisation test was not reported in one study, and another did not report if tests were conducted on non-HD or HD days or were standardised for follow-up.	<p>Report:</p> <ul style="list-style-type: none"> <li>- Timing of testing in relation to HD treatment;</li> <li>- Familiarisation procedures;</li> <li>- Procedures for collecting patient reported measures;</li> <li>- Explicit reporting of standardisation of these procedures for follow-up.</li> </ul>
	Poor reporting of adverse events	Five trials reported whether any adverse events had occurred. Statements about adverse events lacked detail.	<ul style="list-style-type: none"> <li>- Adhere to CONSORT reporting guidance, including specific guidance for reporting harms(69);</li> <li>- Use of standardized and validated measurement instruments for adverse events, where possible.</li> </ul>
	Reporting bias in relation to study results	Selective outcome reporting in five of studies due to incomplete reporting of outcomes or time-points, reporting of only statistically significant outcomes. Only one study had published a protocol paper.	<ul style="list-style-type: none"> <li>- Registration of trials;</li> <li>- Publication of protocol paper;</li> <li>- Presentation of all data in numerical form, not solely figures;</li> <li>- Adherence to Consort for NPT(54);</li> <li>- Raw data freely available in repository.</li> </ul>
<b>R e d</b>	Attrition bias	Of the ten studies providing information about	Reduce dropout:

		attrition, a mean of 23% (range 6-56%) of participants were lost to follow-up in the IDC group, and 14%, (range 0-56%) in usual care. Only one study reported the use of an intention to treat analysis for their primary outcome measure.	<ul style="list-style-type: none"> <li>- Minimise number of visits or integrate into usual care;</li> <li>- Allow step-wise withdrawal if unavoidable;</li> <li>- Engage a patient involvement group to provide patient perspective on trial procedures;</li> <li>- Judicious selection of outcome measures;</li> <li>- Feasibility studies prior to full scale trial.</li> </ul> <p>Undertake an intention to treat analysis. Explicitly state the methods used to address missing data.</p>
	Small sample sizes and underpowered studies	<p>Only three studies based their sample sizes on an a priori power calculation for their primary outcome. One was unable to recruit the required number of patients.</p> <p>Trials of exercise and rehabilitation can be challenging to deliver and the numbers of patients required to adequately power some outcomes may be large depending on the chosen primary outcome measure</p>	<ul style="list-style-type: none"> <li>- A priori power calculation for primary outcome measure for RCTs or report as a feasibility or pilot study, with appropriate outcomes, limitations and conclusions;</li> <li>- Collaborative working to increase sample sizes and facilitate multi-centre working (key for some outcomes e.g., echocardiogram).</li> </ul>
	Lack of blinding of participants and intervention providers	Three studies reported that they were unable to blind participants and personnel, others do not report blinding.	<ul style="list-style-type: none"> <li>- Report blinding explicitly including methods and the fidelity;</li> <li>- Where blinding not possible, strategies to reduce bias should be reported;</li> </ul>
	Lack of blinding of outcome assessors	Blinding of outcome assessment was reported incompletely in only five studies, all of which were high risk due to explicit lack of blinding or researchers assisting participants to complete QoL questionnaires.	<ul style="list-style-type: none"> <li>- Aim to use blinded outcome assessors where possible;</li> <li>- Report blinding explicitly including methods and the fidelity;</li> <li>- Where blinding not possible, strategies to reduce bias should be reported.</li> </ul>
	Baseline imbalances	Five studies demonstrated large, potentially	<ul style="list-style-type: none"> <li>- Adequate sample sizes and randomisation</li> </ul>

		clinically significant differences between control and IDC groups for baseline LVMI, SBP, the SF-36 and some measures of physical function, although all authors state that these differences were not statistically significant.	<p>procedures;</p> <ul style="list-style-type: none"> <li>- Identification of important baseline differences a priori;</li> <li>- Discuss characteristics of important differences in results;</li> <li>- Do not use significance testing to determine if baseline imbalances exist.</li> </ul>
<b>Enhanced synthesis of trial results</b>	Wide heterogeneity of outcome measures	Limited number of outcomes appropriate to combine in meta-analyses.	<ul style="list-style-type: none"> <li>- Use of core outcomes as identified by SONG-HD in addition to outcomes pertinent to the aims of the specific trial (66);</li> <li>- Outcomes selected are appropriate to trial design;</li> <li>- Validation/ reliability studies of these functional measures for the population.</li> </ul>
	Discuss clinical significance of results.	Individual trials report statistically significant results that seem unlikely to translate to a meaningful benefit for patients.	<ul style="list-style-type: none"> <li>- Report confidence intervals and effect sizes;</li> <li>- Comment on the clinical relevance of findings within the results alongside statistical significance;</li> <li>- Use MCIDs from other chronic disease populations where ones for HD do not exist.</li> </ul>

613 **Table 2: Recommendations for enhancing the reporting, quality and synthesis of clinical trials of IDC. (CONSORT, CONSolidated**  
614 **Standards of Reporting Trials; HD, haemodialysis; IDC, intradialytic cycling; LVMI, left ventricular mass index; MCID, minimum**  
615 **clinically important difference; NPT, non-pharmacological trials; QoL, quality of life; RCT, randomised controlled trial; SBP, systolic**  
616 **blood pressure; SF-36, Medical Outcomes Short Form 36; SONG-HD, Standardised Outcomes in Nephrology- haemodialysis; TIDieR,**  
617 **Template for intervention description and replication.).**

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620 **Legends to figures**

621 Figure 1: Flow diagram of study selection. \*: some studies excluded for multiple reasons.

622 (HD, haemodialysis; IDC, intradialytic cycling, RCT, randomised controlled trial).

623

624 Figure 2: Forest plot comparing IDC with usual care on VO<sub>2</sub> peak (A) and the six-minute

625 walk test (B).

626

627 Figure 3: Forest plots comparing change in physical (A) and mental component summary

628 scores of the SF-36 (B) in IDC and usual care.

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630 Figure 4: Forest plot comparing IDC with usual care on systolic (A), diastolic blood pressure

631 (B) and pulse wave velocity (C).

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