ABSTRACT

Objectives: For people with end-stage renal disease requiring haemodialysis, exercise can improve aspects of quality of life (QoL). However, the relative benefits and risks of different types of exercise in this population are unknown. Therefore, this pilot study aimed to evaluate the feasibility of a main study evaluating the efficacy of cycling and resistance exercise each performed during the haemodialysis treatment on QoL.

Methods: In this factorial (2×2) pilot trial, 31 haemodialysis patients were randomised to cycling, resistance, cycling and resistance, or an attention control. Feasibility was defined a priori by criteria on recruitment, fidelity to the protocol and patient response to the intervention. To better understand feasibility, we conducted interviews with dialysis unit staff and trial participants. As secondary outcomes, we estimated the main effect of cycling and weights each compared with control on QoL, physical function and strength.

Findings: We exceeded the target accrual of 28 participants over 12 weeks. Irrespective of exercise group allocation, adherence was high; of the 1038 training sessions offered, 87% were initiated and over 80% of exercise sessions were performed as per protocol. Progression based on perceived exertion, individual instruction and interactions with the kinesiologist facilitated acceptability across exercise groups. Using an attention control, measures of contamination and attrition were low. Important barriers to unit staff readiness for the intervention were initial safety and workflow concerns, unit workload and onerous data collection. Secondary outcomes were not statistically significant. Adverse events were low and did not increase with a higher volume of exercise.

Conclusions: The main study is feasible with minor modifications. In addition to practical assistance, involvement from unit staff could increase patient participation and improve trial implementation. Strategies to increase acceptability of the intervention for staff include improving workflow integration and using a pretest demonstration phase to introduce the intervention.

Trial registration number: NCT02234232. Results

INTRODUCTION

While haemodialysis (HD) is a life-sustaining therapy for people with end-stage renal disease (ESRD), it is associated with low quality of life (QoL) and a marked decline in functional status. Although the benefits of exercise in this population have been recognised, few studies have evaluated how different types of exercise can influence QoL, and the majority of interventions have evaluated aerobic exercise. How to most effectively engage patients in the optimal exercise prescription and achieve the desired outcome while minimising risk, is critical to increasing patient participation.

Many generic QoL scales used in exercise studies in people with ESRD address the individual’s perception of their ability to meet the demands of everyday living. However,
The performance of daily tasks is more dependent on musculoskeletal fitness than aerobic capacity. In the elderly non-ESRD population and in people with congestive heart failure (CHF), resistance training is a promising means of improving QoL and decreasing disability. However, whether resistance training confers specific benefits relevant to aspects of QoL in people with ESRD is not known.

The aim of our future multicentre study is to evaluate the effect of two types of exercise (cycling and resistance) each compared with control and performed during the HD treatment (intradialytic exercise, IDE) on QoL and physical performance using a randomised factorial design. Prior to proceeding with this main study, a pilot was warranted to evaluate the feasibility of the design. Although delivering exercise during HD has been associated with greater adherence compared with a home-based exercise programme, few pilot studies have rigorously evaluated the feasibility or the integrity of trial implementation and we are not aware of any studies that have included qualitative methods to provide a more comprehensive understanding of the implementation process.

**METHODS**

**Study design**

This mixed methods, single-centre, randomised, factorial (2×2) trial included qualitative interviews with trial participants and dialysis unit staff to evaluate domains of feasibility defined a priori: recruitment, fidelity to the study protocol and the response of trial participants and dialysis unit staff to the intervention. In a secondary analysis, we explored differences in QoL, physical function and strength. The two factors evaluated were aerobic exercise (cycling) and resistance exercise (leg weights). HD patients were randomised to one of four groups: cycling, leg weights, combined leg weights and cycling, or stretching (an attention control). The rationale for using a factorial design is for the efficiency of testing more than one intervention in the same participants. (There is no known interaction between aerobic and resistance exercise in the literature i.e., the effect of aerobic exercise does not differ in the presence of resistance exercise.) All exercises were performed during HD at each thrice-weekly dialysis session over 12 weeks (36 sessions). The study protocol was registered under NCT02234292.

**Setting and participants**

The trial setting was an outpatient dialysis unit in Edmonton, Canada that serves ~110 patients. A study coordinator recruited participants during their HD sessions. Inclusion criteria were as follows: adult (age ≥18); dialysis dependent for ≥3 consecutive months; receiving ≥3 dialysis treatments per week; mobile (any distance, walking aid permitted); at least one non-prosthetic limb; and capable of providing consent. Exclusion criteria were as follows: currently enrolled in a clinical trial; missing an average of >2 dialysis sessions per month; planned move or modality change within the next 4 months; currently enrolled in a structured exercise programme; scheduled hospitalisation for >1 week; unstable during HD; and any uncontrolled medical condition that would preclude participation in a low/moderate intensity exercise programme.

**Randomisation and blinding**

Participants were randomised on a 1:1:1:1 ratio using a computerised randomisation procedure with permuted blocks of eight and twelve. Allocation was concealed in serially numbered, opaque, sealed envelopes. The randomisation list was generated by the statistician and kept in a locked cabinet. Given the open setting of the dialysis unit and the nature of the intervention, it was not logistically possible to blind the participants or the kinesiologist to treatment allocation. Therefore, participants and HD unit staff were blinded to the study hypothesis. Patients were informed that they would be randomised to one of four different exercise regimens; a stretching exercise group served as the attention control. Kinesiologists assessed all tests of physical performance; a blinded assessor performed outcome assessments at 12 weeks.

**Exercise intervention**

A kinesiologist instructed all participants on how to perform exercises and supervised a minimum of two of the participants’ thrice-weekly exercise sessions. In addition, the kinesiologist supervised the first three exercise sessions and the first session following progression of the exercise prescription. When the kinesiologist was not present, dialysis unit staff assisted patients with equipment set-up and completed trial documentation. Throughout the study, unit staff were also asked to help motivate patients by providing verbal encouragement. The kinesiologist instructed all participants on how to use rating of perceived exertion (RPE) with the Borg scale (6–20). The intensity of exercise for the aerobic, resistance and combined intervention groups was prescribed at a level of 12–14 or ‘somewhat hard’ on the Borg (RPE) scale and a RPE level of 8–9 (‘very light’) for the stretching group.

**Aerobic intervention**

Each session included a 5 min warm-up and cool-down on the cycle ergometer at an RPE of 9–11. The cycling protocol started with 15 min of cycling with time increased by 2.5 min each week. The resistance was adjusted to maintain the target RPE. One of two types of cycle ergometers were used according to compatibility with the type of dialysis chair: the Monark 881E cycle (Healthcare International, Langley, Washington, USA) or the TherapyTrainer (Interactive Motivation, Greeley, Colorado, USA).
Resistance intervention

Ankle weights (Fabrication Enterprises, White Plains, New York, USA) were used for knee extension, knee flexion and hip flexion. A Thera-Band (Hygenic Corporation, Akron, Ohio, USA) was used for hip abduction. Each session included a warm-up of one set of the four exercises against gravity. Based on RPE, exercises progressed from one set of 10–15 repetitions up to three sets. Weight or resistance was increased when the patient’s RPE was less than target.

Combined intervention

Participants in the combined training group performed the full resistance exercise programme followed by the complete cycling programme.

Attention control

To equalise the effect of cointerventions, the control group performed a non-progressive stretching routine during dialysis. Participants performed two sets, each of four exercises as follows: pelvic tilts, gluteal stretch, calf and hamstring stretch. A Thera-Band Stretch Strap (Hygenic Corporation, Akron, Ohio, USA) was used for the calf and hamstring stretches.

Data collection

Clinical data were collected at baseline via interviews with participants and chart review. Survey data, questionnaires and tests of physical performance were performed at baseline and at 12 weeks. At each session, the following data were recorded on exercise data collection forms (DCFs): pre-exercise and postexercise blood glucose (for diabetics), heart rate (HR), blood pressure (BP), reason for exercise non-participation and early termination, if applicable. During exercise, HR, BP and RPE were documented every 5 min. Data on adverse events (AEs) were collected via interview at each exercise session with the kinesiologist and by chart review.

Primary outcomes

The primary outcome of feasibility was defined by a priori criteria (table 1) and focused on the following: recruitment (rate of accrual, reason for non-participation); fidelity to the protocol (dropout, adherence); response to the intervention (physical activity level outside of the dialysis unit, adoption of the other group’s exercise (contamination)) and acceptability of the intervention.

Recruitment

Previous IDE trials report 20–46% of screened patients were randomised. We estimated that ~85% of the 110 patients in this unit would be available for screening and targeted recruiting 28 participants. Based on the assumption that interested patients may already have preferences concerning exercise that would make randomisation undesirable, unwillingness to be randomised to exercise type was selected as a feasibility criterion. Reason for non-participation in the trial was based on self-report.

Fidelity to the protocol

Based on dropout rates from exercise randomised control trials (RCTs) in people with chronic kidney disease, we defined a high dropout as ≥25% of the study population. Any participant who left the study at any time prior to completing the 12-week exercise programme was defined as a dropout. Adherence was measured to assess patients’ willingness to participate in IDE and to ascertain if the exercises were performed as per protocol (table 1).

Response to the intervention

Acceptability of the exercises was defined as ≥50% of participants reporting that they would like to continue their current IDE programme after the trial is over. The change in physical activity performed outside of dialysis time was measured by self-reported questionnaire and using the Human Activity Profile (HAP). To evaluate whether any participants adopted the other group’s intervention (contamination) outside of dialysis time, patients’ completed questionnaires on the types of activities performed in their leisure time at baseline at 12 weeks.

Qualitative interviews

Detailed information on participants and data collection methods can be found elsewhere. To evaluate barriers to IDE implementation and to inform the content of staff in servicing, we interviewed dialysis unit staff 3 months prior to the start of the trial. To better understand the feasibility of unit staff participation in the delivery of the trial, unit staff members were also interviewed 4 months into the 6-month trial. Unit staff were eligible to participate if the RCT directly affected their workflow and if they had worked in the unit during the trial. Interviews with RCT participants were conducted post-trial participation. All RCT participants were eligible if they were capable of sharing their experiences. Interviews were semistructured with open-ended questions followed by specific prompts on aspects of feasibility. All interviews were audiotaped and transcribed verbatim. For this analysis, interviews were coded using predetermined categories corresponding to our areas of feasibility and analysed to yield a descriptive summary of study findings.

Secondary outcomes

Secondary outcomes were as follows: QoL (the physical component summary (PCS) and the mental component summary (MCS)); tests of physical performance (short physical performance battery (SPPB), 30 s sit-to-stand test and 6 min walk test (6MWT)); an objective measure of strength; and AEs. Testing was carried out at baseline and at 12 weeks, pre-HD on their scheduled HD day.
Quality of life
Participants completed The Kidney Disease Quality of Life Short Form 36 (KDQOL-SF 36). Item scores range from 0 to 100, with higher scores being more favourable. For this pilot, only the mean difference in PCS and MCS are reported.

Tests of physical performance
We used a range of tests to measure physical performance of the lower extremities. The SPPB includes strength (five chair stands), endurance (4 m gait speed) and balance (side by side, semitandem and tandem). Each component is scored from 0 to 4 and is summed SPPB scores between 0 (poor) and 12 (best) performance. The 6MWT was used as a measure of aerobic capacity (distance walked reported in meters) and was performed according to recommendations from the American Thoracic Society. To avoid a ceiling effect and to test muscle endurance, the number of complete getting up and sitting down repetitions performed in 30 s (30 s sit-to-stand (STS 30 s)) was also tested. Muscle strength was measured with the one repetition maximum (1-RM) test using a bilateral leg extension machine for the quadriceps.

Table 1 A priori feasibility criteria and outcomes

| Feasibility criteria | Feasibility outcome |
|---------------------|---------------------|
| Recruitment         | A total of 31 participants over 12 weeks |
|                     | No patients reported randomisation to exercise type as a reason for non-participation. |
| Fidelity to the protocol | A total of 16% of participants dropped out: Cycling n=1, transplanted |
|                     | Resistance n=1, injury from motor vehicle collision |
|                     | Combined n=1, moved dialysis unit |
|                     | Attention control n=2, nausea and vomiting; did not like exercise |
| Adherence (willingness of participants to participate): of all exercise sessions offered,* ≥70% were initiated | A total of 87% of prescribed exercise sessions were initiated: |
|                     | Cycling 89% |
|                     | Weights 83% |
|                     | Combined 90% |
|                     | Attention control 86% |
| Adherence (accordance with the exercise prescription): of all exercise sessions offered, ≥70% were performed at the prescribed time/volume and intensity | A total of 86% of prescribed exercise sessions were performed as prescribed: |
|                     | Cycling 87% |
|                     | Weights 84% |
|                     | Combined 88% |
|                     | Attention control 86% |
| Impact of the intervention | A total of 63% of participants said they would continue with their current exercise |
| Acceptability of the exercises: overall ≥50% of participants reporting that they would like to continue their current intradialytic exercise programme after the study is over | Cycling 50% |
|                     | Weights 50% |
|                     | Combined 100% |
|                     | Stretching 38% |
| Change in the amount of physical activity performed overall: difference in the HAP scores between baseline and 12 weeks† | MAS: |
|                     | Cycling vs no cycling 4.3 (−2.8, 11.5) p=0.3 |
|                     | Weights vs no weights −1.2 (−8.4, 6.0) p=0.7 |
| AAS:                         | Cycling vs no cycling 1.1 (−7.7, 9.9) p=0.8 |
|                     | Weights vs no weights −0.9 (−9.7, 7.8) p=0.7 |
| Difference in the proportion of participants who reported never exercising outside of HD time | Baseline: 39% of participants exercised almost never or never exercising vs 12 weeks: 29% of participants exercised almost never or never (p=0.55) |
| Contamination: any participant who adopted the exercise(s) of another intervention group during the study period | No participants from the cycling, weights or stretching groups reported performing the other group’s exercise |

*Offered sessions exclude sessions lost to study dropout.
†Analysis performed for main effects adjusting for the baseline score and other factor.
AAS, adjusted activity score; HAP, Human Activity Profile; HD, haemodialysis; MAS, maximal activity score.

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Adverse events

AEs were defined a priori and categorised as serious (death, cardiac event, hospitalisation, disability or any life-threatening event) or other (musculoskeletal injury, hypoglycaemia, hypotension, hypertensive urgency (>200 mm Hg systolic or 110 mm Hg diastolic), loss of consciousness, dialysis access complications or any intervention by HD unit staff beyond minimal ultrafiltration). The primary analysis of AEs compared the frequency of events during the exercise session by randomisation group. In a sensitivity analysis, all events occurring during the 12-week intervention period was planned. In both analyses, only the first event per individual was counted (for each type of AE).

Statistical analysis

We summarised baseline data using percentages, medians and IQR, or mean±SD. For secondary outcomes, we explored the effect of aerobic and resistance exercise on QoL and tests of physical performance using the absolute change in score at 12 weeks relative to baseline. To attain the efficiency of the factorial design, all participants who received the aerobic intervention (cycling and the combined group) were compared with all those who did not (resistance and control exercise group) and a similar approach was used for the resistance training group. Analysis of covariance (ANCOVA) was used to adjust for the baseline score and the other intervention (main effect term). Analysis of variance (ANOVA) was performed using Stata Statistical Software, V.13 MP software (http://www.stata.com).

RESULTS

This trial is reported according to the Consolidated Standards of Reporting Trials (CONSORT) guidelines and the recommendations for good practice for the design and analysis of pilot studies.30

Participant flow

Of the 100 patients screened for eligibility, 36 did not meet inclusion criteria and 33 declined to participate (figure 1). The most common reason for exclusion was inability to provide consent (n=8) and the most common reason for declining participation was ‘no interest in exercising during dialysis’ (n=11). A total of 31 participants were randomised and 26 completed the study (cycling, n=7; resistance training, n=6; combined cycling and resistance training, n=7; stretching n=6). Complete outcome data were available for 27 participants.

Baseline characteristics for RCT participants are shown in table 2. Participants were predominantly male (77%), Caucasian (61%), with a median age of 57.5 years (IQR 49.2–75.1). The primary cause of ESRD was glomerulonephritis (32.3%) followed by diabetes (22.6%). In total, 48% of participants were diabetic, 90% had hypertension, 26% had coronary artery disease and 45% of trial participants were taking a β-blocker. Overall, baseline physical functioning was low (mean PCS score of 35±8) and 39% of trial participants reported that they never exercised during their leisure time. Out of 31 RCT participants, 25 participated in interviews (2 declined, 1 had a language barrier and 3 changed location or dialysis modality).

The median age of patient interview participants was 57.5 years (IQR, (IQR) 49.2–68.0); participants were primarily male (76%) and Caucasian (64%). Seven dialysis unit staff participated in pretrial interviews (2 licensed practical nurses (LPNs), 2 registered nurses (RNs), 2 service workers and 1 technician); 86% were female. During the trial, 11 dialysis unit staff were interviewed (2 LPNs, 8 RNs and 1 technician); 91% were female. Two dialysis unit staff participated in both sets of interviews.

Feasibility

Feasibility outcomes are shown in table 1. To highlight key themes regarding the trial’s feasibility, exemplar quotes from the interviews of staff members and patients are shown in boxes 1–4.

Dialysis unit staff (pretrial interviews): barriers to implementation and in servicing

Although none of the staff members who were interviewed had received any prior formal education on IDE, most staff were not interested in attending an educational session. The preferred means of obtaining more information on IDE were by reviewing ‘scientific data’ in their own time. Several staff preferred a practical approach to in servicing and suggested that we focus on teaching them how to set up the exercise equipment and complete study documentation (box 1).

All staff members described potential benefits of IDE, such as improved dialysis and leg cramps, weight loss, increased confidence and patients ‘keeping busy’. However, it was common for staff to express concern that for many patients in the unit, IDE would be unsafe or would interfere with aspects of the dialysis treatment (box 1). Several staff also expressed concern that the exercise equipment would have a negative impact on their workspace.

Dialysis unit staff (pretrial interviews): selection of suitable candidates

Several staff emphasised the importance of selecting appropriate patients for IDE, typically referring to those patients who were stable during HD or younger. Several staff members requested that prior to enrolling a patient, we discuss the patient’s suitability for the trial with them (box 1).
Figure 1  RCT participant flow.

Table 2  Baseline characteristics of trial participants

|                      | All (n=31) | Cycling (n=8) | Weights (n=7) | Combined (n=8) | Stretching (n=8) |
|----------------------|------------|---------------|---------------|----------------|-----------------|
| Age*                 | 57.6 (49.2–75.1) | 66.9 (55.8–82.4) | 59.7 (45.9–81.4) | 60.3 (54.7–68.4) | 49.3 (43.0–62.3) |
| Sex (male)           | 24 (77)   | 8 (100)       | 6 (86)        | 3 (38)         | 7 (88)          |
| Time on HD (years)   | 3.2 (1.7–4.4) | 3.7 (2.4–4.6)   | 2.8 (2.0–4.0) | 2.9 (0.7–2.3) | 3.3 (1.2–6.2) |
| Ethnicity            |            |               |               |                |                 |
| Caucasian            | 19 (61)   | 7 (88)        | 3 (43)        | 5 (63)         | 4 (50)          |
| Southeast Asian      | 4 (13)    | 1 (13)        | 1 (14)        | 1 (13)         | 2 (25)          |
| Aboriginal           | 3 (10)    | 0             | 2 (29)        | 0              | 1 (13)          |
| Other                | 5 (16)    | 0             | 1 (14)        | 2 (25)         | 1 (13)          |
| Cause of ESRD        |            |               |               |                |                 |
| Diabetes             | 7 (22.6)  | 2 (25)        | 1 (14.3)      | 2 (25)         | 2 (25)          |
| Glomerulonephritis   | 10 (32.3) | 1 (12.5)      | 5 (71.4)      | 4 (50)         | 0               |
| Hypertension         | 1 (3.2)   | 1 (12.5)      | 0             | 0              | 0               |
| Polycystic kidney disease | 3 (9.7) | 1 (12.5)      | 0             | 1 (12.5)       | 1 (12.5)        |
| Reflux/urological    | 3 (9.7)   | 1 (12.5)      | 0             | 0              | 2 (25)          |
| Other                | 5 (16.1)  | 2 (25)        | 1 (14.3)      | 0              | 2 (25)          |
| Unknown              | 2 (6.5)   | 0             | 0             | 1 (12.5)       | 1 (12.5)        |
| BMI                  | 24.7 (21.6–29.9) | 23.6 (22.2–25.7) | 25.9 (24.6–29.9) | 25.3 (20.0–30.8) | 24.2 (20.4–33.8) |
| Diabetes             | 15 (48)   | 3 (38)        | 3 (43)        | 5 (63)         | 4 (50)          |
| Hypertension         | 28 (90)   | 8 (100)       | 7 (100)       | 7 (88)         | 6 (75)          |
| β blocker            | 14 (45)   | 4 (50)        | 4 (57)        | 3 (38)         | 3 (38)          |
| Coronary artery disease | 8 (26) | 4 (50)        | 1 (14)        | 2 (25)         | 1 (13)          |
| Heart failure        | 7 (23)    | 4 (50)        | 3 (43)        | 0              | 0               |
| QoL-PCS              | 35±8      | 35±9          | 32±9          | 35±10          | 36±3            |
| Never exercise in leisure time | 12 (39) | 3 (38)        | 4 (57)        | 1 (13)         | 4 (50)          |

*Median (IQR interval); N with (%) or mean (±SD); totals do not always add to 100 due to rounding.
HD, haemodialysis; BMI, body mass index; ESRD, end-stage renal disease; QoL-PCS, quality of life-physical component summary.
Patients' decision to participate in IDE
Several staff stated that patients' social networks in the unit were an effective means of disseminating information. Another staff member stated that after being approached for study participation, patients commonly elicited their opinion (box 1).

Based on the data from the pretrial interviews, modifications were made to the study protocol (table 4).

RCT participants: recruitment
We exceeded the target accrual of 28 participants over 12 weeks. Randomisation to exercise intervention was not a barrier to participation. Patient interview participants reported that recruitment posters displayed outside of the unit and hearing other participants discuss their participation in the trial were effective means of promoting interest and participation in the study (box 2).

Dialysis unit staff (midtrial interviews): fidelity to the protocol
Although the physical demand of delivering the exercise equipment to patients was not described as onerous, data collection for the trial was. One staff stated that there were occasions when trial documentation 'didn't get done'. Several staff reported that there...
were technical challenges with retrieving HR and BP data for DCFs from the HD machines. Some staff also mentioned that recording the vital signs was too time-consuming.

Unit staff frequently made reference to the study as ‘just one more thing’ and trial resource material was not frequently accessed. Although some staff members felt prepared to assist with the trial, several staff suggested that a lack of clarity on trial processes was a barrier to their involvement (box 4).

**RCT participants: fidelity to the protocol**

The dropout rate over the study period was lower than our prespecified threshold at 16%. Irrespective of exercise group allocation, patients’ willingness to participate in IDE and their adherence to the exercise prescription was high; of the 1038 training sessions offered, 87% of sessions were initiated (89% in the cycling group, 83% in the weights group, 90% in the combined group and 86% in the stretching group). The exercises were performed as per protocol within all four groups for >80% of exercise sessions (table 1). Exercise parameters are shown in table 3. For the active intervention groups, the mean RPE was within the targeted range and BP and HR followed a similar trend: increasing during exercise and returning towards baseline post exercise. For the attention control, HR and BP were unchanged over the exercise period.

Although the exercises were protocolised, many participants viewed the intervention as tailored to their level

| Table 3 Exercise parameters for the four exercise groups |
|-----------------|-----------------|-----------------|-----------------|
| **Cycling**     | **Weights**     | **Combined**    | **Stretching/control** |
| Borg (Intensity, RPE) | 13±1            | 13±1            | 13±1            | 8±2            |
| Mean amount of exercise performed | 28.0±3.4 min | 36±12 (repetitions) | 27.5±8.8 min; 35±12 (repetitions) | NAP |
| Systolic BP (mm Hg) | Pre: 136±20       | Pre: 123±26       | Pre: 121±28       | Pre: 119±22       |
|                   | During: 150±26     | During: 127±27     | During: 126±24     | During: 119±22     |
| Diastolic BP (mm Hg) | Pre: 75±16         | Pre: 117±26        | Post: 116±26       | Post 118±20       |
| Heart rate (bpm) | Pre: 66±14         | Pre: 66±15         | Pre: 62±13         | Pre: 70±14         |
|                   | During: 85±20       | During: 63±15       | During: 67±13       | During 70±15       |
|                   | Post 77±17          | Post 74±13          | Post 73±11          | Post 77±17       |

BP, blood pressure; lbs, pounds; RPE, rating of perceived exertion.

Pre, post, and during exercise BP and HRs are a means±SD for initiated exercise sessions.

| Table 4 Modifications to the study protocol following pretrial interviews with unit staff |
|-----------------|-----------------|-----------------|
| **Trial protocol item** | **Initial plan/barrier** | **Modification** |
| In-servicing format | Didactic sessions on the benefits of exercise in people with ESRD and one practical session with the exercise equipment | Two practical in-services on study procedures and equipment set-up Video posted on YouTube on the exercises and how to assist patients with equipment set-up Education materials (articles, pamphlets, summaries) on IDE placed on the unit for staff |
| Workspace safety for staff | Exercise equipment as workspace hazard | Unit staff identified where equipment would be stored on the unit With unit staff input, protocols for equipment set-up and removal were written into study protocol |
| Recruitment | Only study staff selects suitable candidates | Prior to enrolling a patient, the charge nurse was consulted regarding any dialysis-related safety concerns |
| Implementation | Include several unit staff members as volunteer ‘exercise champions’ to lead unit staff and liaise with study staff | No volunteers found. Identified four staff ‘point people’ who were already in leadership roles in the unit to informally check in with study staff on trial implementation |

ESRD, end-stage renal disease; IDE, intradialytic exercise.
Thompson S, et al. BMJ Open 2016;6:e012085. doi:10.1136/bmjopen-2016-012085

Box 4 Exemplar quotes from interview participants on fidelity to the protocol

**Dialysis unit staff (mid-trial interviews)**

“…we check patients every half-hour for their blood pressures and all the dialysis machine readings and stuff like that, so I find also recording the blood pressure is very time-consuming, because we can go back and look at the list of blood pressures on their machine after, but then we just go back and find them or you have to be recording them every 5 or so minutes, so you’re running back and forth between doing your other work and so forth. So I find it’s very busy in that respect…”

‘It was just difficult to add something for us to do, ’cause initially, I think what the thought was to teach all the nurses what the patients were supposed to be doing, but it was just difficult to in-service everybody. They were, like, “Okay, so this is how you fill out the sheet”—’cause the sheet, to me, I’m so confused working with it. And sometimes—oftentimes, we’re short-staffed, so we don’t have the staffing to even get this equipment and all that kind of stuff. So it ended up being they just ended up coming every run and doing the exercise study with the patients. … I think there was a lot of resistance from staff to really help out with it.’

‘I am prepared because they also have an in-service, and they also have [the kinesiologist] here to show us, she also give us e-mail and show with the video, show us how the exercise going. But I be honest, we don’t have time to look at that. We don’t have time to sit down and look at that video!’

**RCT participants (post-trial participation)**

“Yes, because I was starting from zero exercise, so I wasn’t sure how much, how hard it would get, how—if I could keep up to what they wanted, that kind of thing…But they did it very gradual, and [the kinesiologist] was very good about telling us ahead of time when they’re going to put up the weights or when they’re going to increase the minutes of pedaling, so you knew what to expect.’

“Well, we were increased at our own pace, which I really liked, because I just went at my own level.’

‘Also I want to tell you that I have a treadmill at home, but sometimes I do it, sometimes I don’t. But here, it’s, like, we have to...’

 Deus ex Machina (box 4). Individualised instruction, progression based on RPE and support from the kinesiologist21 were commonly mentioned as strengths to the exercise programme. For several patients, knowing there was the expectation of having to exercise facilitated adherence (box 4).

Of the 1038 exercise sessions that were offered, only three were terminated early. In all exercise groups, the most common reason for not initiating a given session was a physical symptom (7.5% of all prescribed sessions), commonly fatigue or feeling generally unwell. HD-related issues accounted for only 1% of non-initiated sessions, primarily due to central venous catheter dysfunction. Many patients mentioned that consistently obtaining exercise equipment from unit staff was the main barrier to exercise participation;21 however, this reason for non-participation was not captured with the exercise DCFs. Only 1.5% of DCFs had missing data for reason not initiated.

Dialysis unit staff (mid-trial interviews): impact of the intervention

Overall, dialysis unit staff agreed that the exercise programme was valuable for patients (box 3). Their perception of benefit was based on patient report, as the trial results were not known at the time of their interviews. Staff viewed patients’ subjective improvements, such as ‘feeling healthier’, as valid evidence of the benefits of IDE (box 3).

RCT participants: impact of the intervention

Across all exercise groups, the patients’ response to exercise was highly favourable (box 3); 92% of participants reported they wanted to continue IDE after the trial and 63% wanted to continue exercising with their current regimen (table 1). There were no crossovers during the trial and no change in the amount of physical activity performed outside of HD time was detected. Concealment of stretching as an active treatment was successful among patients and staff. One participant in the attention control withdrew from the study because he did not find stretching beneficial, ‘it wasn’t straining, it was just too easy’. Although another participant stated that stretching was ‘boring’, most participants in the control group viewed stretching as an important aspect of an exercise regimen (box 3). One participant commented that their exercise routine was shorter than the other groups resulting in relatively less interaction time with the kinesiologist.

Patients commonly discussed the benefits of IDE, and for many, these results motivated them to continue exercising (box 3). Patients discussed the exercise-related benefits of IDE, such as greater strength and endurance. Several patients attributed improvements in daily functioning to participation in IDE. Improvements in dialysis-related symptoms were also mentioned, primarily decreased cramping and restless legs. The most frequently discussed benefit of IDE was that it ‘helped kill the four hours’ and that it made the time on dialysis more enjoyable. For one participant, IDE served as ‘an escape from the humdrum’.

Secondary outcomes

The absolute differences in scores for secondary outcomes are shown in table 5. Scores are presented as crude mean differences and main effects. No significant differences from baseline to 12 weeks were found in the PCS or MCS components of the SF-36 or physical performance tests (6MWT, STS 30 s, 1-RM). For the main effects analysis of the SPPB, the absolute difference in score and (95% CI) was 1.7 (0.2 to 3.3) for the of all those allocated to receive cycling (cycling plus both interventions) versus no cycling (weights plus the attention control) and 1.6 (0.05 to 3.2) for the main effect of those allocated to receive weights versus no weights. This result is consistent with a minimal clinically important difference (values from 0.5 to 1.3 have been recommended).23 31 Interaction terms for the planned
primary outcomes of interest for the main study were as follows: PCS −4.2 (−16.1 to 7.6); p=0.47 and SPPB −2.9 (−5.5 to −0.38); p=0.026.

No serious AEs were reported during the exercise sessions. Owing to the low frequency of events in the trial overall, comparative statistics were not performed. AEs occurring during exercise are shown in figure 2. Two patients in the combined group had AEs (one dialysis access complication, one episode of hypertensive urgency and one episode of hypotension). Two patients in the cycling group had AEs (two episodes of hypertension and ankle abrasions from the bike). In the weights group, there was one episode of access complication. There were no AEs during exercise in the stretching group. The overall frequency of AEs was low (figure 3).

**DISCUSSION**

The purpose of this pilot study was to evaluate the feasibility of an IDE exercise intervention and to perform an exploratory analysis of cycling and weight training each compared with control on QoL, tests of physical performance and strength. We demonstrated feasibility of recruitment and high patient acceptability. In addition, few exercise trials in this population have attempted to blind participants to group allocation.4 We demonstrated a low risk of contamination and attrition with the use of an attention control and blinding to study hypothesis. However, primarily based on the findings from the interviews with dialysis unit staff and trial participants, several modifications to the study protocol are required prior to proceeding with the main study.

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**Table 5**  Secondary outcomes (QoL, tests of physical performance and strength)

| Outcome | Cycling (n=8) | Weights (n=7) | Combined (n=8) | Stretching/control (n=8) |
|---------|--------------|--------------|---------------|-------------------------|
| PCS     |              |              |               |                         |
| Mean difference and SD | 5.2±9.3 | 4.1±8.0 | 1.7±7.4 | 3.4±7.3 |
| Main effects (95% CI) | Cycling vs no cycling | −0.076 (−5.9 to 5.8); p=0.979 | Weights vs no weights | −1.82 (−7.7 to 4.1); p=0.53 |
| MCS     |              |              |               |                         |
| Mean difference and SD | −2.3±10.7 | −3.4±9.1 | −1.5±5.9 | 0.70±7.5 |
| Main effects (95% CI) | Cycling vs no cycling | 0.23 (−6.0 to 6.5); p=0.94 | Weights vs no weights | 0.21 (−6.5 to 6.9); p=0.95 |
| SPPB    |              |              |               |                         |
| Mean difference and SD | 1.9±2.4 | 1.4±1.9 | 1.0±1.2 | 0.63±1.2 |
| Main effects (95% CI)* | Cycling vs no cycling | 1.7 (0.2 to 3.3) p=0.028 | Weights vs no weights | 1.6 (0.05 to 3.2) p=0.044 |
| 6MWT    |              |              |               |                         |
| Mean difference and SD | 42.3±88.8 | 54.9±52.9 | 39.0±76.8 | 0.8±44.0 |
| Main effects (95% CI) | Cycling vs no cycling | 12.8 (−36.1 to 61.6) p=0.60 | Weights vs no weights | 30.7 (−17.8 to 79.2) p=0.21 |
| STS 30 s |              |              |               |                         |
| Mean difference and SD | 0.9±2.2 | 1.6±2.7 | 1.4±3.5 | 1.4±4.3 |
| Main effects (95% CI) | Cycling vs no cycling | −0.31 (−2.7 to 2.1) p=0.79 | Weights vs no weights | 0.42 (−2.0 to 2.8) p=0.73 |
| 1-RM    |              |              |               |                         |
| Mean difference and SD | 11.6±10.7 | 8.9±5.5 | 4.9±11.6 | 9.3±10.1 |
| Main effects (95% CI) | Cycling vs no cycling | −3.4 (−11.0 to 4.2) p=0.37 | Weights vs no weights | −2.8 (−9.9 to 4.2) p=0.42 |

1-RM, one repetition maximum; 6MWT, 6 min walk test; MCS, mental component score; PCS, physical component score; QoL, quality of life; SPPB, short physical performance battery; STS 30 s, 30 s sit-to-stand.

Models are adjusted for baseline score and the other main effect term.

*Interaction term included in the model (p=0.026).
Readiness for change is considered critical to the successful implementation of complex interventions in healthcare settings. In this pilot, we found that there was a lack of readiness among dialysis unit staff for IDE. Several of the factors that influenced unit staff’s preparation, motivation and ability to participate in this trial have been cited in other studies as barriers to the implementation of clinical IDE programmes: lack of time, high patient care demands and safety concerns with the exercise equipment in their workspace. In our previous study, we also identified a lack of support from management and personal beliefs about exercise as influencing staff readiness for IDE. Therefore, prior to recruitment for the main study, it will be necessary to develop a strategy for understanding staff readiness at potential study sites. Although the influence of education on staff participation in IDE remains unknown, in one study, patient and staff thought that a better understanding of IDE would have improved their initial participation. In this pilot, the lack of interest among many unit staff for IDE education was a barrier to engaging staff. Other more convenient forms of delivering education, that is, videos online and reading material on the unit were not highly accessed. As unit staff expressed that seeing and hearing the benefits from their patients first-hand positively influenced their perceptions of the intervention, a pretrial demonstration phase may be the most effective means of promoting acceptability of IDE. Despite the concerns expressed in the pretrial interviews about patient and workspace safety, that no unit staff mentioned these concerns in the second set of interviews (once the intervention was established), also supports the value of providing staff with the opportunity to experience IDE in their own setting prior to study start.

In addition to requiring the unit staff’s assistance with IDE delivery for practical reasons, we identified other reasons why their participation was important. First, due to their frequent and prolonged contact with patients, dialysis unit staff are in a unique position to assist patients with decision-making. As we found that some patients seek the opinion of dialysis unit staff on study participation, it is important that those who engage in these discussions are prepared to discuss the risks and benefits of IDE with patients. Although 30% non-participation is comparable to other trials in this population, it is possible that the staff’s perceptions of IDE influenced patients’ decision to participate. Second, the patients’ perspective that unit staff’s assistance and encouragement with IDE is consistent with their role as carer and patient advocate has the potential to influence patient acceptability of IDE. Third, many patients experienced difficulty consistently obtaining exercise equipment from unit staff, which has clear implications for patient adherence.

For unit staff, exercise data collection was too time-consuming and resulted in missing data. This issue was recognised early in the trial and resolved with greater involvement from study staff. This strategy is not feasible for a multisite study and exercise vital signs will be limited to pre-exercise, midexercise and postexercise. We also found that for unit staff, feasibility of workflow integration was affected by the timing of when in the dialysis treatment that IDE was performed. To decrease the risk of hypotension, other trials have typically completed exercise within the first 1–2 hours of the HD session and starting exercise within the first hour of HD is often recommended. However, this is often the busiest time for unit staff, and in settings where there are staffing constraints, it may be a barrier to optimal staff engagement. We are only aware of one trial where IDE was performed in the final 2 hours of the HD session and this was well tolerated. Our protocol specified that patients finish their exercise within the first 3 hours of the dialysis shift. The safety of this approach is supported by our BP and safety data. A more detailed evaluation of the timing of the HD session and its effect on BP would provide important insight into how to optimise the safety and the practicality of IDE delivery.

This study has several important strengths. Most studies evaluating exercise adherence in people with kidney disease have focused on individual determinants and not evaluated programme factors. In this study, progression based on RPE and individualised instruction facilitated acceptability among patients. As described in our qualitative study, patients perceived the kinesiologist’s technical support as conveying a sense of esteem and capability. This interaction may have served to increase participation, irrespective of group assignment. Additionally, the most commonly mentioned benefit to IDE was that it helped pass the time, suggesting that many patients are interested in participating in interventions where they can use their time on HD more constructively. It also suggests that some of the perceived improvement in well-being could be mediated through engagement in an activity, rather than exercise. These findings underscore the importance of continuing to use a supervised attention control for the main study.
Our study also has several limitations that warrant mention. Given the potential impact of the interaction with the exercise specialist on IDE acceptability, it will be important to ensure that the interaction time between the attention control group and the kinesiologist is equivalent to that of the intervention groups. Also, the trial study population was small and relatively homogeneous with respect to sex, age and ethnicity, which may limit the generalisability of the findings.

We did not detect differences in physical activity or exercise performed outside of the unit during the trial, nor was the trial powered for this outcome. The antagonistic interaction term for the SPPB will also need to be explored in more detail, as this could be a spurious finding due to multiple outcome testing. The primary aim of this pilot study was to evaluate feasibility and small sample sizes were used. Therefore, the finding that cycling or weights did not improve QoL or other measures of physical performance should not be interpreted as providing evidence for no effect. Based on 80% power to detect a difference in the primary outcome of PCS of five points in the main effect of aerobic and the main effect of resistance, 32 participants per arm are required. Allowing for 25% dropout per arm, the main study will enrol 160 patients. A four-arm parallel design would allow direct comparisons between the interventions; however, the sample size would need to be at least twice as large as that calculated for the main study. Given that recruitment and retention are barriers to performing adequately powered exercise studies in this population, the factorial design is one means of improving efficiency while allowing for indirect comparisons between aerobic and resistance training.

CONCLUSIONS

To our knowledge, this is the first feasibility study to use qualitative methods to evaluate IDE implementation within an RCT design and to address known limitations to trial design. In addition to informing the design of our future definitive study, these results are useful in the development of future trials and for guiding clinicians with the implementation of their own IDE interventions. The key lesson learnt was that within this protocolised setting, the potential for unit staff readiness to influence aspects of feasibility, such as recruitment and patient adherence was high. Therefore, prior to study start, more time will need to be invested in understanding and enhancing staff readiness. For engaging unit staff, a less didactic approach that is also integrated into their existing workflow may be highly effective.

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Contributors Authorship followed International Committee of Medical Journal Editors (ICMJE) guidelines. ST was responsible for the conception and design of the project and prepared the manuscript. MH, SK, AM and MT participated in the design of the study and provided methodological input. IG participated in the design of the exercise intervention. AL provided statistical support. All authors read and approved the manuscript.

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