RESEARCH

Exploratory insights into novel prehabilitative neuromuscular exercise-conditioning in total knee arthroplasty

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Abstract

Background: Contemporary strategies for prehabilitation and rehabilitation associated with total knee arthroplasty (TKA) surgery have focused on improving joint range-of-motion and function with less emphasis on neuromuscular performance beneficially affecting joint stability. Furthermore, prehabilitation protocols have been found to be too long and generic-in-effect to be considered suitable for routine clinical practice.

Methods: A pragmatic exploratory controlled trial was designed to investigate the efficacy of a novel, acute prehabilitative neuromuscular exercise-conditioning (APNEC) in patients electing TKA. Adults electing unilateral TKA were assessed and randomly allocated to exercise-conditioning (APNEC, n = 15) and usual care (Control, n = 14) from a specialised orthopaedic hospital, in the United Kingdom. APNEC prescribed nine stressful exercise-conditioning sessions for the knee extensors of the surgery leg, accrued over one week (3 sessions·week−1; 36 exercise repetitions in total; machine, gravity-loaded) and directly compared with usual care (no exercise). Prescribed exercise stress ranged between 60%—100% of participant’s daily voluntary strength capacity, encompassing purposefully brief muscular activations (≤1.5 s). Baseline and follow-up indices of neuromuscular performance focusing on muscle activation capacity (electromechanical delay [EMD], rate of force development [RFD] and peak force [PF]) were measured ipsilaterally using dynamometry and concomitant surface electromyography (m. rectus femoris [RF] and m. vastus lateralis [VL]).

Results: Group mean ipsilateral knee extensor muscular activation capacity (EMDREF [F(3,57) = 53.5; p < 0.001]; EMDVL [F(3,57) = 50.0; p < 0.001]; RFD [F(3,57) = 10.5; p < 0.001]) and strength (PF [F(3,57) = 16.4; p < 0.001]) were significantly increased following APNEC (Cohen’s d, 0.5—1.8; 15% to 36% vs. baseline), but unchanged following no exercise control (per protocol, group by time interaction, factorial ANOVA, with repeated measures), with significant retention of gains at 1-week follow-up (p < 0.001).

Conclusions: The exploratory APNEC protocol elicited significant and clinically-relevant improvement and its retention in neuromuscular performance in patients awaiting TKA.

Trial registration: (date and number): clinicaltrial.gov: NCT03113032 (4/04/2017) and ISRCTN75779521 (3/5/2017).

Keywords: Pre-surgery training, Total knee arthroplasty, Joint replacement, Sensorimotor, Neuromuscular, Exercise-conditioning

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Background

Many recipients of total knee arthroplasty (TKA) are content with the outcome of surgery as an optimal solution for pain relief compared to physical rehabilitation.
Exercise conditioning’s rationale is both intuitively and pragmatically appealing because preoperative enhancements to functional and fitness capabilities should lead to patients experiencing better surgical outcomes and reduced risks of further injury [1] and greater risk of falls [2, 3]. Amongst heterogeneous preoperative exercise methodologies and levels of evidential quality within meta-analytical reviews, significant improvements (small to moderate effects sizes) have been observed in postoperative function, quadriceps strength, and length of stay in patients undergoing TKA [4]. Similarly, while preoperative progressive resistance training (PRT) has been shown to be safe and not liable to exacerbate knee joint pain and effusion [5], the methodological quality of evidence had rendered the effectiveness of PRT before and/or after TKA as inconclusive [6]. Nevertheless, other reports support the use of short-term (4-week) high-intensity resistance training before TKA [7] and capable of inducing long-lasting effects on muscle strength [8], but not necessarily on functional performance [8] or other patient-reported outcomes [7]. Pronounced gains in functional, neuromuscular performance capabilities and higher patient satisfaction levels [9, 10] in related orthopaedic applications involving innovative and physiologically-principled resistance conditioning indicate the potential for improvement on contemporary approaches in which purposely-designed programmes can still have clear benefits for patients awaiting TKA.

Thus, while there is emerging evidence for the efficacy of preoperative resistance training in contemporary literature for improving strength performance, we would argue that the exercise protocols that have been tested thus far [7, 8, 11, 12] may not have not been optimal with regard to their characteristics of physiological stress. As such, we hypothesise that a new approach incorporating ideals for neuromuscular conditioning may elicit more benefits.

Recent exploratory work in asymptomatic adults has shown efficacy within condensed, physiologically-principled approaches to conditioning [13]. As such, promoting short durations of muscle activation’ patterns during exercise in order to minimise the likelihood of pain nociceptor’ activation (<1.5 s [14]) and unwanted provocation of autogenic and arthrogenic inhibition may be beneficial. Short durations of exercise conditioning may offer benefits in terms of efficacy, patients’ burdens and their adherence to prescribed programmes of exercise.

This paper reports on the exploratory findings of ipsilateral responses to acute prehabilitative neuromuscular exercise-conditioning (APNEC) in order to characterise its efficacy in patients awaiting TKA. It is part of a larger study investigating conditioning effects on both ipsilateral and contralateral limbs. Our hypothesis was that an acute and short dosage (one week, intermittently) of physiologically-principled exercise-conditioning, featuring high intensity (60% to 100% of daily strength capacity) and short durations of muscular activation, would show efficacy for improving neuromuscular performance. This study aimed to identify the efficacy of APNEC for improving neuromuscular performance capacities in patients awaiting TKA. Additionally, the retention of effects was explored at one week following cessation of APNEC.

Methods

Study design and participants

A UK single-centre NHS randomised controlled trial was registered (clinicaltrial.gov: NCT03113032 and ISRCTN75779521) and given Research & Development (RIAH/RL1 715) and ethical approval (South East Scotland Research Ethics Committee 01 [IRAS 198,930; REC reference 17/SS/0005]). The trial received endorsement by means of routine patient and public involvement meetings at the Robert Jones and Agnes Hunt (RJAH) Orthopaedic NHS Foundation Trust. Inclusion criteria involved patients over 18 years, diagnosed with severe OA of the knee and undergoing primary TKA surgery. Patients suffering from rheumatic or ongoing neurological disorders, conditions affecting function such as amputation, reduced mental capacity affecting abilities to follow prescribed exercises, or undergoing TKA for diseases other than OA, were excluded. Patients recording previous joint replacement surgery (contralateral knee, or hip) were not excluded.

The screening of the patients was undertaken within the hospital’s outpatient department by the clinicians having oversight (PG; AB). Inclusion criteria involved patients over 18 years, diagnosed with severe OA of the knee and undergoing primary TKA surgery. Patients gave written informed consent prior to baseline assessment (T1). Allocation to two groups (block randomisation [n, 14]) underwent experimental intervention to the ipsilateral leg, receiving nine, 20-min focal exercise-sessions delivered on alternate days (3 sessions per week and 3 sessions per day) during a single week, with clinical oversight (AMR). The Control group (n = 14) followed current practice (no exercise).

Assessment procedures and data capture

Participants were assessed ipsilaterally on four separate occasions that coincided with health care appointments...
for patients: T1 (baseline; ≈ 11 weeks prior to surgery, at initial clinical consultation, including familiarisation with assessment procedures and apparatus), T2 (pre-intervention; two weeks pre-surgery, at appointment for prophylactic infection control), T3 (48 h post-intervention; one week pre-surgery, immediate APNEC effects) and T4 (one week after cessation of intervention; retention of effects at TKA surgery).

Acute prehabilitative neuromuscular exercise-conditioning (APNEC)

APNEC prescribed nine stressful exercise-conditioning sessions for the ipsilateral knee extensors, accrued over one week (3 exercise days [24 h recovery]; 3 sessions per day [typically, 10.00 am; 1.00 pm; 4.00 pm]; 1 exercise per session; 4 repetitions per exercise, with 20 s inter-repetition rest periods [15] [36 repetitions in total] on a calibrated machine, gravity-loaded [Life Fitness, model number FZLE-500023; www.lifefitness.com] (Fig. 1a) and directly compared with usual care (no exercise). Prescribed exercise stress ranged between 60%—100% of participant’s daily voluntary strength capacity (one-repetition maximal [1RM], assessed using 2 or 3, 2-s concentric contractions and 15 min prior to first daily APNEC session) (Table 1), encompassing purposefully brief muscular activations (~1.5 s). APNEC sessions were preceded by standardised warm-up (walk: ≈ 150 m, waiting area to gym) and recovery (240 s [16]), with an exercise repetition’ movements practised twice using an unloaded lever-arm (Fig. 1b, c, d and e), prior to exercise. Physiological stimuli for adaptation were delivered by micro-cyclical patterning of exercise intensity, recovery and progression, featuring high to low intra-day exercise intensity amongst daily sessions. Physiological stimuli for adaptation were delivered by micro-cyclical patterning of exercise intensity, recovery and progression,

Fig. 1 Knee extensor machine and settings a side view b knee position 1–10; this also helped secure the 90° starting position of the knee. c front view d padded leg rest setting: Small, Medium, Large and Extra large; this position determined where the padded leg rest will place pressure on the shin and was normally placed above the ankle e back seat rest setting 1–10; this was normally positioned so that the back of the knee rested comfortably over the machine’s seat-edge, with the shin hanging perpendicularly to the ground.
featuring high to low intra-day exercise intensity amongst daily sessions. The clinical model for APNEC had been derived and adapted from pilot research work involving micro- and meso-cyclical training theory and practices amongst elite professional soccer [17]. Figure 2 describes step-by-step, the performance of a single APNEC exercise repetition.

Concentric muscle actions (patient self-paced) fully extended ipsi- and contralateral knees against the machine’s gravity-loaded lever-arm (Fig. 2a, start position). Bilateral full knee extension (Fig. 2b) momentarily preceded voluntary self-paced ipsilateral knee flexion to 45°, away from the lever-arm with maintained full contralateral knee extension (Fig. 2c). Contralateral knee extensor relaxation prompted the lever-arm’s unresisted gravitational loading (Fig. 2d). Ipsilateral extensor’s eccentric muscle action attempted to arrest and counteract the lever-arm’s downward trajectory very briefly (∼1.5 s) at ≈45° (Fig. 2e). A repetition was completed by the involved musculature’s relaxation and concomitant load-dropping to its starting position (Fig. 2f).

### Table 1

An example of the micro-cyclical loading used during the APNEC delivery for participants randomised to this group. The nine sessions (S1—S9) reflect loading-progression delivered over a 1-week period (T2 to T3), with rest days interspersed amongst conditioning days.

| Session (S) | Monday (%MVC) | Tuesday | Wednesday (%MVC) | Thursday | Friday (%MVC) |
|-------------|---------------|---------|------------------|----------|---------------|
| S1          | 65%           | Rest    | S4               | 70%      | Rest          |
| S2          | 100%          | Day     | S5               | 90%      | Day           |
| S3          | 85%           |        | S6               | 80%      | S7            |
| S4          |               |        | S7               | 60%      |               |
| S5          |               |        | S8               | 80%      |               |
| S6          |               |        |                  | 70%      |               |
| S7          |               |        |                  |          |               |
| S8          |               |        |                  |          |               |
| S9          |               |        |                  |          |               |

**Fig. 2** APNEC exercise movement. Please see main text for full explanation.
Assessment of neuromuscular performance

Neuromuscular performance was assessed ipsilaterally focusing on muscle activation capacity (electromechanical delay [EMD]; rate of force development [RFD]) and knee extension strength (peak force [PF]) using dynamometry and concomitant surface electromyography (m. rectus femoris [RF] and m. vastus lateralis [VL]). Participants were seated on a custom-built dynamometer, incorporating a load cell (range 3 kN; Tedea-Huntleigh, Cardiff, UK) [18, 19]. Knee and hip joint angles were standardised at 45° (0° representing full extension) [20] and 110° [21], respectively. Electromyographic (EMG) recordings were obtained using bipolar rectangular surface electrodes (self-adhesive, Ag/AgCl; 10 mm diameter; Unilet, UK), applied longitudinally and parallel to the muscle fibres’ orientation (30 mm inter-electrode distance [9, 18]; < 5 kΩ impedance [9, 22, 23]). EMG signals were pre-amplified (1902 Mk IV; Cambridge Electronic Design, UK; input impedance 10,000 MΩ, CMRR 100 dB; < 5 kΩ). Knee and hip joint angles were standardised at 45° (0° representing full extension) [20] and 110° [21], respectively. Electromyographic (EMG) recordings were obtained using bipolar rectangular surface electrodes (self-adhesive, Ag/AgCl; 10 mm diameter; Unilet, UK), applied longitudinally and parallel to the muscle fibres’ orientation (30 mm inter-electrode distance [9, 18]; < 5 kΩ impedance [9, 22, 23]). EMG signals were pre-amplified (1902 Mk IV; Cambridge Electronic Design, UK; input impedance 10,000 MΩ, CMRR 100 dB; < 5 kΩ). EMG recordings were obtained using bipolar rectangular surface electrodes (self-adhesive, Ag/AgCl; 10 mm diameter; Unilet, UK), applied longitudinally and parallel to the muscle fibres’ orientation (30 mm inter-electrode distance [9, 18]; < 5 kΩ impedance [9, 22, 23]). EMG signals were pre-amplified (1902 Mk IV; Cambridge Electronic Design, UK; input impedance 10,000 MΩ, CMRR 100 dB; < 5 kΩ).

Participants received an assessor’s verbal signal to activate (as rapidly and forcefully as possible) and relax their knee extensors (after 2 s to 3 s of activation), undertaking three static contractions separated by 10-s rest periods. Interrogation of raw EMG and concomitant force–time records (exceeding 95% confidence limits of the background electrical noise amplitude) [18, 19], identified peak performance of EMDRF and VL (shortest latency [ms]), PF (highest force response [N]) and RFD (maximum rate at which force is increased (N s⁻¹) amongst the three trials. Prior to the assessment, participants warmed-up via subjectively-judged static contractions (~2 s; 10 s, inter-repetition neuromuscular recovery [15]) at 25% (2 repetitions), 50%, 75% and 100% of maximum voluntary contraction [18, 19].

Statistical analysis

Group means (± SD) described outcome scores. Separate factorial (group [APNEC; Control] x time [T1; T2; T3; T4]) analyses of variance (ANOVAs), with repeated measures for time, tested hypotheses relating to EMDRF, EMDVL, RFD and PF using per protocol analyses (SPSS Vn. 23, IBM SPSS Illinois, USA). A priori reverse Helmert orthogonal difference testing located anticipated time-specific effects. Any violations of assumptions underpinning the use of ANOVA (for example, sampling from a normally distributed population, sphericity) were countered using Greenhouse–Geisser (GGE) adjustments. Statistical significance was accepted at p < 0.05, with Cohen’s d quantifying relative effect size (ES: 0.2, 0.5 and 0.8 considered in general to be ‘small’, ‘medium’ and ‘large’ effects, respectively) and percent change relative to ‘baseline’ performance offered as an additional marker of change in performance. A priori experimental design sensitivity estimation offered an approximate statistical power of 0.7 for avoiding intrusion of type-II errors for a medium relative effect size (Cohen’s d; 0.5) in the study’s primary outcome, EMD, and at its end-point (T4), requiring an approximate sample size of n = 12 within each group (www.sportsci.org).

Results

Twenty-nine patients from 238 candidates electing unilateral TKA (standardised implant procedure [MRK™]) on orthopaedic waiting-lists (May 2017 to April 2018) participated. Statistical analyses were undertaken on data from the 21 patients who completed the study protocol (Fig. 3: CONSORT enrolment; Table 2: Per protocol participants’ characteristics).

Neuromuscular performance

Factorial interactions (group x time) showed group mean ipsilateral knee extensor muscular activation capacity (EMDRF [F(3,57) = 53.5; p < 0.0005]; EMDVL [F(3,57) = 50.0; p < 0.001]; and RFD [F(3,57) = 10.5; p < 0.001]) and strength (PF [F(3,57) = 16.4; p < 0.001]) were significantly increased immediately following APNEC, but unaffected by no exercise control (Table 3). Performance improvements between baseline and immediately after APNEC (EMDRF [Cohen’s d, 0.62; 25.5%; F(1,19) = 53.5; p < 0.0005, a priori difference contrast]; EMDVL [d, 0.66; 26.0%; F(1,19) = 50.0; p < 0.0005]; RFD [d, 0.54; 13.2%; F(1,19) = 10.5; p < 0.001]; PF [d, 0.55; 18.7%; F(1,19) = 16.4; p < 0.001]) were prominent, contributing most to the overall ANOVA interactions, and were retained substantively at one week after APNEC’s cessation (T4: EMDRF [Cohen’s d, 0.87; 14.7% vs. baseline; F(1,19) = 26.5; p < 0.001, a priori difference contrast]; EMDVL [d, 0.80; 15.0%; F(1,19) = 26.8; p < 0.001]; RFD [d, 0.39; 9.5%; F(1,19) = 8.2; p < 0.001]; PF [d, 0.46; 16.1%; F(1,19) = 7.4; p < 0.001]) (Fig. 4).

Discussion

This exploration study of prehabilitative neuromuscular exercise-conditioning in patients electing unilateral TKA showed the protocol’s efficacy for improving neuromuscular performance capacities, with substantial gains in peak ipsilateral knee extensor muscular activation characteristics (EMD [Cohen’s d, 0.62 – 0.66; 25.5 – 26.0% and RFD [d, 0.54; 13.2%]) and strength (PF [d, 0.55; 18.7%]) following APNEC. Concomitant responses of participants acting as controls were trivial (d < 0.2; < 3.0%; ns) and reflected contemporary practice in which no structured exercise was undertaken and only variations in activities of daily living might have elicited serendipitous conditioning stimuli.
Optimum dose–response characteristics for APNEC await scrutiny, but this formulation involving a short period of dosage, high intensity and purposely brief nociceptor response-evading muscular actions, elicited responses exceeding statistical, precision and reliability criteria for the selected indices of neuromuscular performance (≈ 4%—8%) [19, 20] and appears to offer important clinical relevance in counteracting persistent performance deficits [24, 25].

Relevant comparable potency data amongst the contemporary literature is elusive for all outcomes. Nevertheless, peak strength gains following APNEC (PF [d,
Table 3  Group mean scores (± SD) for per protocol assessments at baseline (mean T1, T2), T3 and T4 for the leg undergoing surgery in the APNEC (n = 9, for all time points of assessment [5 males; 4 females]) and Control (n = 12, for all time points of assessment [7 males; 5 females]) groups. Calculated effect sizes (Cohen’s d) and percentage changes are relative to baseline performance

| Outcome | Baseline | T3 | T4 | ES Baseline – T3 | ES Baseline – T4 |
|---------|----------|----|----|-----------------|-----------------|
| EMDRF (ms) |           |    |    |                 |                 |
| APNEC   | 43.5 ± 7.2 | 33.5 ± 6.0 | 38.4 ± 7.1 | 1.78 | 0.96 |
| Control | 45.0 ± 7.7 | 43.6 ± 7.5 | 45.3 ± 7.1 | 0.20 | 0.02 |
| EMDVL (ms) |           |    |    |                 |                 |
| APNEC   | 46.3 ± 8.1 | 33.9 ± 6.4 | 38.9 ± 8.3 | 1.70 | 0.91 |
| Control | 46.8 ± 8.1 | 45.3 ± 7.6 | 46.8 ± 7.4 | 0.20 | 0.01 |
| RFD (N·s⁻¹) |        |    |    |                 |                 |
| APNEC   | 57.8 ± 127.2 | 658.7 ± 143.5 | 637.2 ± 145.7 | 0.59 | 0.43 |
| Control | 605.0 ± 114.8 | 587.2 ± 128.5 | 610.5 ± 134.0 | 0.10 | 0.01 |
| PF (N)   |           |    |    |                 |                 |
| APNEC   | 182.4 ± 55.1 | 211.9 ± 62.8 | 207.1 ± 66.3 | 0.50 | 0.40 |
| Control | 171.8 ± 42.8 | 171.7 ± 44.9 | 172.8 ± 43.7 | 0.01 | 0.00 |

Key: EMD electromechanical delay (ms), RF rectus femoris muscle, VL vastus lateralis muscle, RFD rate of force development (N·s⁻¹), PF peak force (N)

0.55]) matched or exceeded those amongst studies of established rehabilitative conditioning in patients undergoing TKA surgery [26, 27] or following conservative treatment [28, 29] (d ≤ 0.50). APNEC’s capability for improving several independent facets of neuromuscular performance (correlations amongst EMD, RFD and PF, at baseline: r < 0.40; ns) could be deployed usefully as a strategic alternative to aspects of contemporary prehabilitative practices associated with TKA, within end-stage OA in general, or serve as a specific augmentation. In the absence of definitive minimal clinically-important difference criteria [30], it was notable that gains for all APNEC patients exceeded the minimum detectable change criteria for EMD (13.8 ms; 19) and the performance changes of all control patients (Fig. 5). Retention of gains in peak ipsilateral knee extensor muscular activation characteristics (EMDFS [Cohen’s d, 0.91 – 0.96; 53.5%, proportion of gain as a percentage]; EMDVL [Cohen’s d, 0.96; 53.9%] RFD [d, 0.43; 72.8%]) and strength (PF [d, 0.41; 80.0%]) following APNEC’s cessation were substantial, significantly better than baseline performance scores (p < 0.05) and had similarly exceeded the relevant minimum detectable change criteria.

There is emerging evidence for the efficacy of pre-operative resistance training in the contemporary literature for improving strength performance, including some protocols using relatively short durations (4 weeks) [7, 8, 11, 12]. By contrast, the efficacy of APNEC was derived from a more notably truncated protocol involving just nine stressful exercise-conditioning sessions for the ipsilateral knee extensors, accrued over one week, but encompassing deliberately brief muscular activations.

The APNEC protocol’s short duration emphasises its potential for utility within clinical time-logistics. It also favours neural mechanisms rather than morphological adaptation [31] underpinning the gains in capabilities. There is neuromuscular performance capabilities. There is potential for APNEC’s characteristically brief, eccentric muscular actions to provoke favourable changes in neural activation patterns [32], with increased rates of motor unit firing [28]. Autogenic and arthrogenic inhibition may be reduced consequently by increased rates of motor firing, provoking sympathetic concomitant gains in neuromuscular performance [29, 32]. The complementarity of APNEC’s brief but intense muscle actions in mitigating nociceptor activation and potentiating increased activity amongst previously inhibited motor units may be crucial for efficacy in this clinical population. Purposely sequenced delivery of episodes of APNEC might feasibly offer potentiated effects on neuromuscular performance capacities in this clinical population, which could counter clinical criticisms of contemporary prehabilitative exercise programmes and their ability to elicit only subtle and transient improvements [33]. Future well-controlled clinical trials should continue to explore optimal modes and dose-responses of APNEC delivery in this population, which may occur in the absence of competing physiological conditioning stimuli, such as concomitant strength- and endurance-related exercise stresses [9].

Limitations to this study were related to its delivery and design. Logistical and ethical constraints had precluded assessments in the immediate period after surgery, and evaluation of longer-term decay patterning of APNEC gains in neuromuscular performance amongst complex
interactive effects associated with surgery and post-TKA clinical care. Similarly, experimental controls were focused on an extended period of longitudinal evaluation of baseline performance capabilities of the leg undergoing surgery and differential inter-group responses, rather than on those of the contralateral leg.

Other limitations for this explorative trial included that group allocation was not concealed from patients or from those overseeing the patients’ testing administration and conditioning. As such, the potential for bias within the findings could not be eliminated. Similarly, physical activity behaviors associated with travel to and from the APNEC’s venue for its delivery and assessments was not monitored directly and varied physical activity might have elicited heterogeneous carry-over effects amongst the patients’ responses to APNEC. A further limitation reflected a lack of patients’ self-perceived pain assessments within the APNEC protocol, even though the latter had been monitored in general by means of questionnaire, but not reported here. Furthermore, while the patient’s compliance with the APNEC’s training prescription was monitored directly, this approach

Fig. 4 Group mean performance scores (± SD; APNEC, n = 9; Control, n = 12) assessed in the ipsilateral leg for muscle activation capacity (electromechanical delay [EMD]; rate of force development [RFD]) and knee extension strength (peak force [PF]) using dynamometry and concomitant surface electromyography (m. rectus femoris [RF] and m. vastus lateralis [VL]).
that may not be facilitated in all environments, such as within self-managed care. Nevertheless, future studies will aim to identify optimised APNEC dosing and approaches for its scalability and delivery amongst varied care environments.

This study’s findings were derived from a modestly-sized sample of non-obese (BMI < 30 kg·m⁻²) patients (n = 21), which might preclude generalisation. Observed Type II error rates were modest (≤0.12) and appeared to offer suitable experimental design sensitivity and statistical power amongst the selected indices of neuromuscular performance.

**Conclusion**

This exploration study of prehabilitative neuromuscular exercise-conditioning in patients electing unilateral TKA suggested that the APNEC protocol may be efficacious for improving neuromuscular performance capacities (d, 0.54—0.66; 13.6—26.0%). Gains prevailed the cessation of APNEC.

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**Authors’ contributions**
All authors contributed extensively to the work presented in this paper. AMR, MvdL, AB, PG and NG participated in the conception and design of the manuscript. AMR, MvdL, AB and NG contributed to the drafting of the manuscript and conducted data collection. NG and MvdL supervised the study, with clinical oversight by AB and PG, and AMR and NG analysed the data. All the authors discussed the results and study procedures, contributing extensively to the work presented in this manuscript, and approved the final version.

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Availability of data and materials
The datasets used and analysed during the current study are available from the corresponding author on reasonable request.

Declarations

Ethics approval and consent to participate
South East Scotland Research Ethics Committee, in conjunction with Research & Development Dept., RIAH Orthopaedic NHS Foundation Trust and Queen Margaret University, Edinburgh Research Ethics Committee. Informed consent was obtained from each participant and all methods were carried out in accordance to the relevant guidelines and regulations.

Consent for publication
Not applicable.

Competing interests
None.

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