A telerehabilitation programme in post-discharge COVID-19 patients (TERECO): a randomised controlled trial

Jian’an Li,1,2 Wenguang Xia,3 Chao Zhan,4 Shouguo Liu,1,2 Zhifei Yin,1 Jiayue Wang,1,2 Yufei Chong,3 Chanjuan Zheng,3 Xiaoming Fang,4 Wei Cheng,4 Jan D Reinhardt

ABSTRACT

Objectives To investigate superiority of a telerehabilitation programme for COVID-19 (TERECO) over no rehabilitation with regard to exercise capacity, lower limb muscle strength (LMS), pulmonary function, health-related quality of life (HRQOL) and dyspnoea.

Design Parallel-group randomised controlled trial with 1:1 block randomisation.

Setting Three major hospitals from Jiangsu and Hubei provinces, China.

Participants 120 formerly hospitalised COVID-19 survivors with remaining dyspnoea complaints were randomised with 61 allocated to control and 59 to TEREKO. 

Intervention Unsupervised home-based 6-week exercise programme comprising breathing control and thoracic expansion, aerobic exercise and LMS exercise, delivered via smartphone, and remotely monitored with heart rate telemetry.

Outcomes Primary outcome was 6 min walking distance (6MWD) in metres. Secondary outcomes were squat time in seconds; pulmonary function assessed by spirometry; HRQOL measured with Short Form Health Survey-12 (SF-12) and mMRC-dyspnoea. Outcomes were assessed at 6 weeks (post-treatment) and 28 weeks (follow-up).

Results Adjusted between-group difference in change in 6MWD was 65.45 m (95% CI 43.8 to 87.1; p<0.001) at post-treatment and 68.62 m (95% CI 46.39 to 90.85; p<0.001) at follow-up. Treatment effects for LMS were 20.12 s (95% CI 12.34 to 27.9; p<0.001) post-treatment and 22.23 s (95% CI 14.24 to 30.21; p<0.001) at follow-up. No group differences were found for lung function except post-treatment maximum voluntary ventilation. Increase in SF-12 physical component was greater in the TEREKO group with treatment effects estimated as 3.79 (95% CI 1.24 to 6.35; p=0.004) at post-treatment and 2.69 (95% CI 0.06 to 5.32; p=0.045) at follow-up.

Conclusions This trial demonstrated superiority of TEREKO over no rehabilitation for 6MWD, LMS, and physical HRQOL.

Trial registration number ChiCTR2000031834.

INTRODUCTION

After discharge from acute care, many survivors of COVID-19 experience ongoing symptoms, impairment of pulmonary function, decreased exercise capacity, reduced muscle strength, activity limitations and reduced quality of life.1-7 Problems may persist for at least 6 months.1 This indicates the need for the provision of rehabilitation services that can decrease the burden on patients and the health system.8 Pulmonary rehabilitation measures with demonstrated effectiveness in COPD9 and, with low-certainty evidence (one trial), SARS10 are obvious candidates.

Evidence from high-quality trials on the effectiveness of such programmes in COVID-19 survivors is, however, lacking to date.11 Moreover, delivery of conventional inpatient or outpatient rehabilitation is complicated through diminished capacity in postacute care as well as clinical and public health measures imposed to reduce the risk of viral transmission.12 Telerehabilitation provides a viable alternative that could be superior to no rehabilitation and as effective as conventional rehabilitation.13 14 We investigated possible superiority of a telerehabilitation programme for COVID-19 (TERECO) over no rehabilitation with regard to functional exercise capacity, lower limb muscle strength...
(LMS), pulmonary function, perceived dyspnoea and health-related quality of life in formerly hospitalised COVID-19 survivors. We further report on the occurrence of adverse events.

**METHODS**

**Study design**

Multicentre, parallel-group randomised controlled trial. The original protocol for this study is available from (URL: http://idmr.scu.edu.cn/info.htm?id=1841614474692833).

**Setting, recruitment and consent**

Three centres from Jiangsu (Jiangsu Province Hospital/Nanjing Medical University First Affiliated Hospital), Hubei Wahan (Hubei Province Hospital of Integrated Chinese and Western Medicine) and Hubei Huangshi (Hubei Huangshi Hospital of Chinese Medicine) recruited patients recovering from COVID-19. In total, 1242 patients had been discharged from these hospitals when we stopped recruitment on 28 May 2020 of which about one-third (n=377) was deemed potentially eligible after prescreening of hospital records. Possible candidates were then contacted by telephone and an appointment for a baseline visit was made. At this baseline visit, further assessment of eligibility was performed and written informed consent was obtained.

**Participants**

Participants were aged 18–75 years, discharged from one of the participating hospitals after inpatient treatment for COVID-19, and had modified British Medical Research Council (mMRC) dyspnoea score of 2–3. The latter inclusion criterion was chosen as we anticipated that patients with moderate remaining dyspnoea symptoms could actively participate in the programme and most benefit from it. Moreover, as this was an unsupervised intervention, patients with mMRC dyspnoea of 4–5 were excluded for reasons of safety. Other exclusion criteria were: resting heart rate over 100 bpm, uncontrolled hypertension, uncontrolled chronic disease (eg, diabetes with random blood glucose >16.7 mmol/L, haemoglobin A1C >7.0%), cerebrovascular disease within 6 months, intra-articular drug injection or surgical treatment of lower extremities within 6 months, taking medication affecting cardiopulmonary function such as bronchodilators or β-blockers, unable to walk independently with assistive device, unable or unwilling to collaborate with assessments, enrolled or participated in other trials within past 3 months, having history of severe cognitive or mental disorder or substance abuse, enrolment in other rehabilitation programme.

**Random sequence generation and allocation concealment**

Permutated allocation sequences for 1:1 block randomisation (block size 10–14) stratified by hospital were computer-generated by an independent statistician. Allocation was concealed by central randomisation and only revealed after baseline assessment through call to study centre.

**Blinding of assessors**

Baseline visits of each potential study participant involved one assessor (rehabilitation doctor) and one independent allocator (therapist). Assessors left the study site after the baseline measurements. Allocators then contacted the study centre in the presence of the patient to reveal allocation. Patients and therapists were requested to not disclose allocation to assessors at any time during the study.

**Procedures**

**Control group**

Participants in the control group received short educational instructions at baseline.

**TERECO group**

Participants took part in an unsupervised 6-week home exercise programme delivered through a smartphone application called RehabApp and monitored with a chest-worn heart rate (HR) telemetry device. Teleconsultations with therapists were carried out once per week. The exercise programme involved 3–4 sessions per week. It included (i) breathing control and thoracic expansion, (ii) aerobic exercise and (iii) LMS exercises specified in a three-tiered exercise plan with difficulty and intensity scheduled to increase over time. Initial exercise types and intensity were determined by physiotherapists contingent on the baseline assessment in accordance with the American College of Sports Medicine’s guidelines for exercise preparticipation. Exercise intensity prescribed for aerobic exercise was based on HR reserve determined by Karvonen’s formula. Intensity ranged from 30%–40% for tier 1 to 40%–60% for tier 3. Having reached at least two-thirds (66.7%) of the scheduled total and effective target time as given in online supplemental table S1 or modified by the therapist in any given week for at least 5 of the 6 weeks was considered compliant with the exercise protocol. Details on interventions and determination of compliance are provided in online supplemental text S1 and table S1.

**Assessments**

Assessments were conducted between 26 April and 9 December 2020. For each patient, home visits were scheduled at baseline, at 6 weeks (post-treatment) and at 24 weeks (follow-up). Additional assessments for dyspnoea and adverse events were performed by consultation via cell phone or WeChat voice call at 2 and 4 weeks. Due to a change of administrative regulations, home visits were no longer permitted for the final follow-up assessment. Participants were thus invited to return to the hospitals where they had originally received treatment. This adjustment led to a delay in the final assessment for about 4 weeks on average.

**Primary outcome**

The primary outcome was functional exercise capacity at post-treatment measured with the 6 min walking test (6MWT) administered in accordance with guidelines from the European Respiratory Society and American Thoracic Society and recorded as 6 min walking distance (6MWD) in metres. For the first two assessment points a course was arranged outside, near the patients’ home. For the final follow-up assessment a course was arranged in the hospital ward according to the same criteria. In each case, the walking course was arranged on flat territory, hard surface, 30 m straight with every 3 m marked by coloured tape and two small cones placed at the turnaround points. Blood pressure was taken before and after the 6MWT and intermittently if indicated. HR and pulse-oxygen saturation were continuously monitored. If patients used walking aids in daily life (eg, rollator, cane), they were allowed to use those during the test. Patients could rest at any time and then continue the 6MWT based on their own assessment and that of the supervising therapist. If the 6MWT could not be completed, distance walked until final interruption was recorded.
Secondary outcomes
The 6MWD at follow-up was assessed as described in the previous section. LMS was measured with the static squat test. This involved a squat against the wall with both feet flat on the ground approximating a 90° angle at hip and knees. Time in seconds participants could remain in squatting position was recorded. Static testing was preferred over dynamic (eg, sit to stand) for ease of standardisation in the home setting. Pulmonary function was evaluated by spirometry according to guidelines of the American Thoracic Society (grade C). A portable pulmonary function device (MINATO, AS-507, Japan) was used. The following parameters were recorded: FEV1 in litres, FVC in litres, FEV1/FVC, maximum voluntary ventilation (MVV) in litres per minute and peak expiratory flow (PEF) in litres per second. Per cent of predicted value for FEV1, FVC and FEV1/ FVC, and per cent below lower limit of normal (LLN) were calculated with Global Lung Initiative 2012 equations for South-East Asia. Per cent of predicted PEF and MVV were calculated with equations for mainland China. Health-related quality of life (HRQOL) was evaluated with the Short Form Health Survey-12 (SF-12). Physical component score (PCS) and mental component score (MCS) are reported, with higher scores indicating better health. Due to the absence of reference equations for mainland China scores were standardised according to US norms. Perceived dyspnoea was assessed with the mMRC scale. Since only patients with moderate dyspnoea (mMRC score of 2 or 3) were included in this study, being dyspnoea-free (mMRC score = 0) was defined as a favourable outcome (coded 1 in analysis). All other mMRC scores were defined as non-favourable outcome (coded 0 in analysis).

Adverse events
Participants could report adverse events at any time during the study via phone call or WeChat message. They were also asked for adverse events during regular assessments. In addition, participants in the TERECO group received a prompt by RehabApp after each session and were asked for adverse events in weekly consultations with therapists. Death, cardiovascular events, other life-threatening events and re-hospitalisation for events related to the intervention or COVID-19 were defined as serious adverse events. All reported adverse events were rated by two independent doctors who had access to event and treatment effect model parameters. Occurrence of a favourable outcome in mMRC-dyspnoea was analysed with a generalised linear model of the Poisson family with a log link, adjusted for centre and using the natural logarithm of the number of observed occasions until the respective data point as offset. Cluster robust SEs (cluster variable: participant ID; number of clusters: 115, average cluster size: 3.9 (min 1, max 4)) were used for the estimation of 95% CIs. Treatment effects are presented as rate ratios (RR) along with logistic 95% CIs. A graphical illustration of trajectories presents estimated marginal means and probabilities (mMRC-dyspnoea) as predicted with the above models by intervention group and time point with 84% CIs (logistic CIs for mMRC) serving as comparisons bars. In contrast to 95% CIs, 84% CIs allow for visual inspection of statistical significance of mean differences at the 5% level by looking at non-overlap of CI bounds for the group means.

Prespecified sensitivity analysis included estimation of the above models on the per-protocol sample, as well as on two types of multiply imputed datasets. First, multiple imputation with chained equations (70 sets) was performed under an extended MAR assumption, that is, that missing values were also dependent on observed values of auxiliary variables not included in aforementioned models (gender, age, disease severity, time from hospital admission to baseline assessment, presence of comorbidities, smoking history, body mass index). Second, controlled multiple imputation (50 sets) was used for simulating a non-MAR scenario where patients with missing assessments in the TERECO group followed the pattern of change in controls (copy increments in reference (CIR)). As a delay in the planned follow-up assessment occurred for several patients and unequal time periods between post-treatment and follow-up assessment resulted, additional post hoc sensitivity analysis was conducted, fitting models with two additional terms for time since onset of symptoms (TOS) and TOS squared.

Analysis of harms
Adverse events were descriptively analysed.

Statistical analysis
All analyses were performed with STATA V.14.0. Main analyses were conducted on intention-to-treat (ITT) basis without imputations. Statistical significance was set at alpha=5% with two-sided tests. Data were assumed to be missing at random (MAR) with missing values depending on observed model parameters. This assumption was tested with sensitivity analysis. The statistical analysis plan (SAP) is available in online supplemental material S2.

The 6MWD was evaluated with constrained longitudinal data analysis, that is a linear mixed effects model that imposed an equality constraint on baseline means. Analogous to analysis of covariance, such equality constraint is imposed to counteract regression to the mean, that is, the statistical phenomenon that a group of patients with worse average baseline scores generally improves more than the group with better scores, independent of a possible intervention effect. The model was adjusted for centre (fixed effect) and dependence of longitudinal observations within study participants was modelled with a random intercept. Parameters for interaction terms between time and treatment represent the estimated treatment effects at post-treatment and follow-up, and are reported with 95% CIs. All secondary outcomes apart from mMRC-dyspnoea were analysed analogously. Occurrence of a favourable outcome in mMRC-dyspnoea was analysed with a generalised linear model of the Poisson family with a log link, adjusted for centre and using the natural logarithm of the number of observed occasions until the respective data point as offset. Cluster robust SEs (cluster variable: participant ID; number of clusters: 115, average cluster size: 3.9 (min 1, max 4)) were used for the estimation of 95% CIs. Treatment effects are presented as rate ratios (RR) along with logistic 95% CIs. A graphical illustration of trajectories presents estimated marginal means and probabilities (mMRC-dyspnoea) as predicted with the above models by intervention group and time point with 84% CIs (logistic CIs for mMRC) serving as comparisons bars. In contrast to 95% CIs, 84% CIs allow for visual inspection of statistical significance of mean differences at the 5% level by looking at non-overlap of CI bounds for the group means.

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Analysis of harms
Adverse events were descriptively analysed.
Figure 1 Flow of patients through the study. ITT, intention to treat; PP, per protocol.

Protocol deviations
Deviations from the study protocol and SAP are described and explained in detail in online supplemental material S3.

Role of the funding source
Funders of the study had no role in study design, data collection, data analysis, data interpretation or writing of this report. The corresponding author had full access to all data in the study and final responsibility for the decision to submit for publication.

RESULTS
Figure 1 illustrates the flow of patients through the study. After prescreening of hospital records, 140 patients were contacted for further evaluation of eligibility between 22 April and May 28 2020. Of those 20 were ineligible or refused consent. One hundred twenty patients were randomised with one person in the TERECO group and one in the control group not receiving the allocated intervention. One patient was withdrawn from the TERECO group before the start of the exercise programme due to premature beat. One patient in the control group had been admitted as a correction of hospital treatment was 26.2 days on average (SD 15.3). Time from hospital discharge to baseline assessment was 70 days on average (SD 16.9). Fifty (43.5%) patients were below LLN for FEV1, 45 (39.1%) for FVC and 26 (22.6%) for FEV1/FVC.

Table 2 gives an overview of crude change and adjusted treatment effects for all outcomes. Figure 2 depicts marginal trajectories over time by study group with 84% CIs (comparison bars) serving as a visual aid for inspecting statistical significance of mean differences at the 5% level (see online supplemental material S5 for detailed estimates).

Primary outcome
The mean 6MWD in the control group increased by 17.1 m (SD 63.9) from baseline to post-treatment assessment, whereas 6MWD in the TERECO group improved by 80.2 m (SD 74.7). The adjusted between-group difference in change in 6MWD from baseline (treatment effect) was 65.45 m (95% CI 43.8 to 87.1; p<0.001).

Secondary outcomes
With estimated 68.62 m (95%CI 46.39 to 90.85; p<0.001), the treatment effect regarding 6MWD increased somewhat at follow-up. LMS improved to a larger degree in the TERECO group as compared with control with estimated treatment effects of 20.12s in squat position (95%CI 12.34 to 27.9; p<0.001) post-treatment, and 22.23s (95%CI 14.24 to 30.21; p<0.001) at follow-up. Lung function parameters improved in both group over time (figure 2). No group differences were found apart from an adjusted between-group difference in change from baseline of 10.57 L/min (95%CI 3.26 to 17.88; p=0.005) in post-treatment MVV in favour of the TERECO group. SF-12 PCS increased to a larger degree in the TERECO group with treatment effects estimated as 3.79 points (95%CI 1.24 to 6.35; p=0.004) at post-treatment and 2.69 (95% CI 0.06 to 5.32; p=0.045) at follow-up. Improvement in SF-12 MCS was somewhat greater in the TERECO group but 95% CIs were also compatible with greater improvement in control at both assessment points. With 90.4% being dyspnoea-free (favourable outcome) in the TERECO group as opposed to 61.7% in the control group (adjusted RR 1.46, 95% CI 1.17 to 1.82; p=0.001), a treatment effect for mMRC-dyspnoea was found immediately after the intervention period but not at the other time points.

Sensitivity analysis
Estimates from sensitivity analysis are provided in table 3. Per-protocol analysis showed larger effect estimates for 6MWD and LMS. For the per-protocol sample the estimated between-group difference in change from baseline for the primary outcome was 72.25 m (95%CI 47.54 to 96.97; p<0.001). With the exception of 6MWD at follow-up, treatment effects were lowest under the CIR scenario, followed by the extended MAR scenario. A long-term effect of TERECO on SF-12 PCS was unstable in all points. Overall, 36 participants in the TERECO group complied with the exercise protocol, 61.02% of those randomised (n=59) and 69.2% of those who remained in the programme for the full 6 weeks (n=52). Compliance increased until week 4, dropped in week 5 and increased again in week 6. More information on missing data and compliance with the exercise protocol is given in online supplemental material S4.

Baseline characteristics of the study participants by intervention group are provided in table 1. The overall mean age of the study population was 50.61 (SD 10.98), 53 (44.5%) were male and 73 (61.3%) had at least one comorbidity. Length of hospital stay for acute treatment was 26.2 days on average (SD 15.3).
Table 1 Continued

| Descriptor | Total (n=119*) | Control (n=60*) | Intervention (n=59*) |
|-----------|---------------|----------------|---------------------|
| FVC below LLN, n (%) | 45 (39.1) | 22 (38.6) | 23 (39.7) |
| FEV1/FVC | 0.80 (0.13) | 0.81 (0.12) | 0.79 (0.14) |
| FEV1/FVC (% predicted), mean (SD) | 96.43 (15.93) | 97.86 (15.03) | 95.03 (16.78) |
| FEV1/FVC below LLN, n (%) | 26 (22.6) | 12 (21.1) | 14 (24.1) |
| MVV (L/min) | 68.72 (28.90) | 63.05 (26.12) | 74.30 (30.60) |
| MVV (% predicted), mean (SD) | 62.69 (22.13) | 58.94 (20.86) | 66.37 (22.88) |
| PEF (L/s) | 3.93 (2.07) | 3.66 (1.75) | 4.21 (2.33) |
| PEF (% predicted), mean (SD) | 48.94 (21.77) | 46.41 (18.20) | 51.42 (24.70) |
| SF-12 PCS, mean (SD) | 39.42 (8.48) | 39.69 (8.25) | 39.15 (8.76) |
| SF-12 MCS, mean (SD) | 44.40 (8.48) | 44.13 (8.25) | 44.67 (8.76) |
| mMRC-dyspnea, n (%) | 2 | 116 (97.5) | 58 (96.7) | 58 (98.3) |
| Other comorbidity | 28 (23.5) | 12 (20.0) | 16 (27.1) |
| Smoking history, n (%) | 15 (12.6) | 6 (10.0) | 9 (15.3) |

During hospitalisation: acute respiratory distress, respiratory rate ≥30 breath/min; pulse oxygen saturation (SpO2) ≤93% at rest; arterial blood partial pressure of oxygen/fraction of inspired oxygen (PaO2/FIO2) ≤300 mm Hg (1 mm Hg =0.133 kPa); respiratory failure requiring mechanical ventilation; septic shock; failure of other organs requiring intensive care unit treatment.46

81.04 (15.20) 80.43 (15.39) 83.62 (14.99)

Continued

preplanned sensitivity analyses. Estimates from post hoc sensitivity analysis that added parameters for TOS to the models were almost identical with those from main analysis.

Adverse events

No serious adverse events occurred during the study period. Eight patients (five in the TERECO and three in the control group) were hospitalised, all for non-life-threatening reasons unrelated to COVID-19 or the intervention and all in the follow-up period. A detailed account of adverse events is provided in the online supplemental material S6 and table S6.1.

DISCUSSION

In this trial, the TERECO programme was superior to no rehabilitation with regard to functional exercise capacity, LMS and physical HRQOL. All these effects could be sustained over a 7-month period. Pronounced differences in exercise capacity and LMS remained between intervention and control group. For physical HRQOL, the difference between TERECO and the control group decreased at follow-up due to improvements in controls. We also found a short-term effect of TERECO on

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Rehabilitation

MVV and mMRC-dyspnoea. Both effects, however, decreased at follow-up with differences no longer being statistically significant. No effects of TERECO on the four other pulmonary function parameters and on mental HRQOL were found. Adherence to the intervention programme was satisfactory and no serious adverse events occurred. While attrition and missing data may have influenced effect sizes, the estimates of exercise capacity and LMS in ITT analysis were consistent with MAR scenarios and more conservative estimates when non-MAR was assumed.

This study evaluated a relatively inexpensive, patient-centred, adaptable telerehabilitation intervention with a wide range of parameters of relevance to function and HRQOL. With a few exceptions, this trial was executed according to the original

### Table 2

| Number of participants | Crude change from baseline* or % endorsing favourable outcome† | Estimated treatment effect (group difference in mean change or risk ratio§) with 95% CI | P value¶ |
|------------------------|-------------------------------------------------------------|--------------------------------------------------------------------------------|---------|
| **Primary outcome**    |                                                             |                                                                                |         |
| 6MWD (m)               |                                                             |                                                                                |         |
| Post-treatment (6 weeks) | 60 | 52 | 17.09±3.94 | 80.20±4.66 | 65.45 (43.80 to 87.10) | <0.001 |
| Follow-up (~28 weeks)  | 55 | 50 | 15.17±7.02 | 84.81±8.38 | 68.62 (46.39 to 90.85) | <0.001 |
| Secondary outcomes     |                                                             |                                                                                |         |
| Squat time (s)         |                                                             |                                                                                |         |
| Post-treatment (6 weeks) | 60 | 52 | 7.98±9.53  | 29.35±27.22 | 20.12 (12.34 to 27.90) | <0.001 |
| Follow-up (~28 weeks)  | 55 | 50 | 4.16±9.62  | 28.12±27.17 | 22.23 (14.24 to 30.21) | <0.001 |
| Pulmonary function     |                                                             |                                                                                |         |
| FEV1 (L)               |                                                             |                                                                                |         |
| Post-treatment (6 weeks) | 56 | 51 | 0.18±0.53  | 0.28±0.51 | 0.00 (−0.18 to 0.17) | 0.969 |
| Follow-up (~28 weeks)  | 53 | 47 | 0.29±0.43  | 0.29±0.48 | 0.00 (−0.18 to 0.17) | 0.969 |
| FVC (L)                |                                                             |                                                                                |         |
| Post-treatment (6 weeks) | 56 | 51 | 0.19±0.40  | 0.21±0.47 | 0.02 (−0.14 to 0.18) | 0.818 |
| Follow-up (~28 weeks)  | 53 | 47 | 0.27±0.43  | 0.30±0.38 | 0.01 (−0.16 to 0.17) | 0.95 |
| FEV1/FVC               |                                                             |                                                                                |         |
| Post-treatment (6 weeks) | 56 | 51 | 0.01±0.16  | 0.04±0.17 | 0.03 (−0.02 to 0.07) | 0.224 |
| Follow-up (~28 weeks)  | 53 | 47 | 0.02±0.15  | 0.02±0.18 | −0.01 (−0.05 to 0.03) | 0.732 |
| MVV (L/min)            |                                                             |                                                                                |         |
| Post-treatment (6 weeks) | 56 | 51 | 5.61±17.31 | 14.49±21.60 | 10.57 (3.26 to 17.88) | 0.005 |
| Follow-up (~28 weeks)  | 53 | 47 | 13.81±20.78 | 18.47±22.31 | 5.20 (−2.33 to 12.73) | 0.176 |
| PEF (L/s)              |                                                             |                                                                                |         |
| Post-treatment (6 weeks) | 56 | 51 | 0.66±1.95  | 0.98±1.90 | 0.38 (−0.24 to 1.00) | 0.229 |
| Follow-up (~28 weeks)  | 53 | 47 | 0.97±1.84  | 0.76±1.92 | −0.02 (−0.66 to 0.62) | 0.954 |
| Quality of life        |                                                             |                                                                                |         |
| SF-12 PCS              |                                                             |                                                                                |         |
| Post-treatment (6 weeks) | 60 | 52 | 3.84±7.60  | 7.81±7.02 | 3.79 (1.24 to 6.35) | 0.004 |
| Follow-up (~28 weeks)  | 55 | 50 | 5.20±9.13  | 8.2±10.05 | 2.69 (0.06 to 5.32) | 0.045 |
| SF-12 MCS              |                                                             |                                                                                |         |
| Post-treatment (6 weeks) | 60 | 52 | 4.17±8.79  | 6.15±10.78 | 2.18 (−0.54 to 4.90) | 0.116 |
| Follow-up (~28 weeks)  | 55 | 50 | 5.51±7.79  | 6.92±10.28 | 1.99 (−0.81 to 4.79) | 0.164 |
| mMRC perceived dyspnoea, to favourable outcome | | | | |
| Interim 2 weeks        | 60 | 54 | 45.0      | 57.4 | 1.27 (0.88 to 1.82) | 0.197 |
| Interim 4 weeks        | 60 | 54 | 61.7      | 66.7 | 1.08 (0.82 to 1.42) | 0.605 |
| Post-treatment (6 weeks) | 60 | 52 | 61.7      | 90.4 | 1.46 (1.17 to 1.82) | 0.001 |
| Follow-up (~28 weeks)  | 55 | 50 | 60.0      | 72.0 | 1.22 (0.92 to 1.61) | 0.162 |

*Crude change from baseline is given as mean change±SD of this change for all outcomes apart from for mMRC (favourable outcome).
† For mMRC, per cent being dyspnoea-free is provided.
‡ Estimated treatment effects for all outcomes apart from mMRC-dyspnoea (favourable outcome) are between-group mean differences in change from baseline derived from mixed effects regression with random intercept for study participant; models are constrained to a common baseline mean across groups and adjusted for centre. Estimation includes all available observations from participants randomised (number of participants with valid observations at baseline is 115 for pulmonary function parameters and 119 for all other outcomes).
§ Estimated treatment effects for mMRC-dyspnoea (favourable outcome) are risk ratios derived from generalised linear model from Poisson family with log link adjusted for centre and ln(number valid observations up to data point) as offset; 95% CIs are based on cluster robust SEs (cluster variable: participant ID).
¶Probability of treatment effect being zero.
CI, confidence interval; MCS, mental component score; mMRC, modified Medical Research Council; MVV, maximum voluntary ventilation; 6MWD, six min walking distance; PCS, physical component score; PEF, peak expiratory flow; SF-12, Short Form Health Survey-12; TERECO, telerehabilitation intervention for COVID-19 survivors.
protocol and attrition was low (about 12%). Sensitivity analysis demonstrated stability of most results under different scenarios.

Limitations of this research include participant characteristics: only COVID-19 survivors with moderate dyspnoea symptoms who had previously been hospitalised for treatment were included. The results are thus not generalisable to persons with mild or severe dyspnoea, nor to people who contracted SARS CoV-2 but were not hospitalised. We further excluded patients taking β-blockers or bronchodilators precluding inference for this population. It should be noted that the intervention might not be suitable for people with very severe impairment and sequelae due to COVID-19 or those not familiar with smartphone technology. Another important weakness is the unexpected change of the location and resulting delay of the final follow-up assessment. It is unclear how this may have affected patient-reported assessments and pulmonary function testing.

Low-certainty evidence (one randomised crossover trial) suggests that 6MWT performed outdoors yields comparable results to centre-based testing. Emerging evidence suggest that the most profound impairment of lung function in COVID-19 occurs in diffusion capacity. This was unclear at the time of study design and the required measurement procedures are difficult to perform at home. SDs of 6MWD used for sample size calculation were based on a study of patients with SARS. This was done for pragmatic reason as no respective data were available for COVID-19 in April 2020. However, although coronaviruses causing both diseases show some degree of genetic similarity, there are also some marked differences in genome and pathology. Future trials may thus refer to the growing body of research in patients with COVID-19. Finally, this trial was not powered for subgroup analysis and effect sizes in specific subpopulations hence remain unclear. For example, length of inpatient stay was longer, and the proportion of patients with severe COVID-19 was somewhat greater in the intervention group which may have enabled a larger effect size.

At the time of writing, there are no other randomised controlled studies on rehabilitation effectiveness for COVID-19. Demonstrating clinically meaningful and sustainable effects of the TERECO programme on 6MWD and LMS, this study adds to previous low-certainty evidence on the effectiveness of telerehabilitation in respiratory disease. The effect size for 6MWD in the present study at 6 weeks is comparable to results from a randomised controlled trial from Hong Kong, which evaluated a 6-week outpatient exercise programme for SARS survivors with baseline and post-treatment assessment. In contrast to our findings, the latter study did not detect any effects of the programme on HRQOL (SF-36) or LMS (measured as gluteus maximum and anterior deltoid strength with dynamometer). While a recent systematic review and meta-analysis reported superior effects of breathing exercise on lung function parameters (FEV1 and FEV1/FVC) as compared with control for COPD, no such effects were found in the present study. A possible explanation is that, in contrast to physical endurance and strength, lung function was not sufficiently targeted by the exercises included in the TERECO programme. This interpretation is supported by our finding that MVV was the only pulmonary parameter that showed a larger increase in TERECO than control at post-treatment. The MVV is not a measure of lung volume but rather of respiratory muscle strength and endurance. Obviously, the latter but not other lung function parameters were targeted by the breathing control and thoracic expansion exercises in the
Rehabilitation

Table 3  Results of sensitivity analysis

| Outcome                       | Estimates of treatment effects from different scenarios with 95% CIs                                                                 |
|-------------------------------|-------------------------------------------------------------------------------------------------------------------------------------|
|                               | ITT, primary analysis (n=119, n_int=59, n_obs=336) | Per protocol‡ (n=91, n_int=36) | ITT, extended MAR multiple imputation* (n=119, n_int=59, 70 sets) | ITT, CIR multiple imputation† (n=119, n_int=59, 50 sets) | ITT, model including time from symptoms onset§ (n=119, n_int=59, n_obs=336) |
| Primary outcome: 6MWD (m)     | 65.45 (43.80 to 87.10) | 72.25 (47.54 to 96.97) | 62.23 (40.07 to 84.39) | 57.18 (35.42 to 78.95) | 65.12 (44.50 to 85.74) |
| Follow-up (−28 weeks)         | 68.62 (46.39 to 90.85) | 75.92 (51.21 to 100.64) | 61.99 (39.22 to 84.76) | 63.07 (40.87 to 85.27) | 67.99 (46.77 to 89.20) |
| Squat time (s)                |                                                                  |                                    |
| Post-treatment (6 weeks)       | 20.12 (12.34 to 27.90) | 22.67 (13.91 to 31.43) | 20.32 (11.72 to 28.91) | 17.81 (10.01 to 25.61) | 20.15 (12.44 to 27.86) |
| Follow-up (−28 weeks)          | 22.23 (14.24 to 30.21) | 25.94 (17.18 to 34.70) | 21.48 (12.73 to 30.24) | 20.07 (12.09 to 28.06) | 22.38 (14.45 to 30.31) |
| Pulmonary function             |                                                                  |                                    |
| FEV1 (L)                      | 0.08 (−0.08 to 0.25) | 0.09 (−0.1 to 0.28) | 0.07 (−0.11 to 0.25) | 0.06 (−0.11 to 0.24) | 0.08 (−0.08 to 0.25) |
| Post-treatment (6 weeks)       | 0.00 (−0.18 to 0.17) | −0.03 (−0.22 to 0.16) | −0.05 (−0.23 to 0.13) | −0.00 (−0.18 to 0.18) | 0.00 (−0.18 to 0.17) |
| FVC (L)                       | 0.02 (−0.14 to 0.18) | 0.09 (−0.08 to 0.27) | −0.01 (−0.18 to 0.16) | 0.03 (−0.13 to 0.20) | 0.02 (−0.14 to 0.18) |
| Post-treatment (6 weeks)       | 0.01 (−0.16 to 0.17) | 0.07 (−0.11 to 0.25) | −0.06 (−0.23 to 0.12) | 0.01 (−0.16 to 0.18) | 0.01 (−0.15 to 0.17) |
| FEV1/FVC                      | 0.03 (−0.02 to 0.07) | 0.02 (−0.02 to 0.07) | 0.02 (−0.02 to 0.06) | 0.02 (−0.02 to 0.06) | 0.02 (−0.02 to 0.06) |
| Post-treatment (6 weeks)       | −0.01 (−0.05 to 0.03) | −0.02 (−0.07 to 0.03) | −0.01 (−0.05 to 0.03) | 0.00 (−0.04 to 0.04) | −0.01 (−0.05 to 0.03) |
| MVV (L/min)                   | 10.57 (3.26 to 17.88) | 14.3 (6.1 to 22.5) | 10.09 (2.11 to 18.07) | 10.32 (2.91 to 17.73) | 10.57 (3.3 to 17.85) |
| Post-treatment (6 weeks)       | 5.20 (−2.33 to 12.73) | 7.29 (−1.00 to 15.59) | 3.04 (−5.38 to 11.46) | 6.01 (−1.77 to 13.78) | 5.25 (−2.27 to 12.76) |
| PEF (L/s)                     | 0.38 (−0.24 to 1.00) | 0.52 (−0.17 to 1.22) | 0.35 (−0.29 to 0.99) | 0.35 (−0.29 to 0.99) | 0.38 (−0.24 to 1.00) |
| Post-treatment (6 weeks)       | −0.02 (−0.66 to 0.62) | −0.16 (−0.86 to 0.55) | −0.18 (−0.83 to 0.48) | 0.03 (−0.62 to 0.67) | −0.03 (−0.67 to 0.61) |
| Quality of life               |                                                                  |                                    |
| SF-12 PCS                     | 3.79 (1.24 to 6.35) | 3.70 (0.76 to 6.63) | 3.68 (1.13 to 6.24) | 3.27 (0.69 to 5.86) | 3.75 (1.22 to 6.27) |
| Post-treatment (6 weeks)       | 2.69 (0.06 to 5.32) | 2.37 (−0.57 to 5.30) | 2.31 (−0.43 to 5.05) | 2.44 (−0.28 to 5.16) | 2.72 (0.12 to 5.33) |
| SF-12 MCS                     | 2.18 (−0.54 to 4.90) | 1.92 (−1.14 to 4.97) | 2.17 (−0.57 to 4.91) | 1.65 (−1.07 to 4.38) | 2.18 (−0.53 to 4.89) |
| Post-treatment (6 weeks)       | 1.99 (−0.81 to 4.79) | 1.48 (−1.58 to 4.54) | 2.30 (−0.48 to 5.09) | 1.82 (−0.96 to 4.61) | 1.93 (−0.67 to 4.73) |
| mMRC dyspnoea, to favourable outcome |                                                                  |                                    |
| Interim 2 weeks               | 1.27 (0.88 to 1.82) | 1.33 (0.90 to 1.98) | 1.26 (0.87 to 1.82) | 1.26 (0.88 to 1.82) | 1.28 (0.89 to 1.85) |
| Interim 4 weeks               | 1.08 (0.82 to 1.42) | 1.04 (0.76 to 1.42) | 1.06 (0.80 to 1.40) | 1.06 (0.80 to 1.40) | 1.09 (0.82 to 1.43) |
| Post-treatment (6 weeks)       | 1.46 (1.17 to 1.82) | 1.43 (1.14 to 1.80) | 1.42 (1.13 to 1.79) | 1.40 (1.11 to 1.77) | 1.47 (1.17 to 1.84) |
| Follow-up (−28 weeks)          | 1.22 (0.92 to 1.61) | 1.15 (0.84 to 1.57) | 1.21 (0.92 to 1.58) | 1.23 (0.92 to 1.63) | 1.23 (0.93 to 1.62) |

Apart from estimates for mMRC perceived dyspnoea (favourable outcome), estimates are between-group differences in mean change from baseline derived from linear mixed effects models adjusted for study centre (fixed effect) and with baseline means constrained to be equal across comparison groups. Estimates for mMRC perceived dyspnoea are rate ratios derived from a general linear mixed model of the Poisson family with log link adjusted for centre and ln(number valid observations up to data point) as offset. CIs are estimated with cluster robust standard errors (cluster variable: participant ID).

* Based on multiple imputation using chained equations assuming data were missing at random. The imputation model included all outcomes and the following auxiliary variables with complete baseline information: gender, age, smoking history (no vs yes), presence of any comorbidity (no vs yes), COVID-19 severity (non-severe vs severe), body mass index, time from admission to hospital to baseline assessment in days.

‡ Based on per-protocol sample (n=119, n_int=59, n_obs=336)

§ Same as main analysis but providing additional adjustment for time from onset of symptoms to measurement point in days and time from onset of symptoms to measurement point in days squared. A likelihood ratio test confirmed superior fit of the model that also included the squared term.

CI, confidence interval; CIR, copy increments from reference; ITT, intention to treat; MAR, missing at random; MCS, mental component score; mMRC, modified Medical Research Council; MVV, maximum voluntary ventilation; 6MWD, six min walking distance; PCS, physical component score; PEF, peak expiratory flow; SF-12, Short Form Health Survey-12.
TERECO programme. Similar to the HRQOL physical component, mental HRQOL also improved in both groups but no statistically significant between-group differences in increments were detected, although the SF-12 MCS score of the TERECHO group remained at about 2 points above control at follow-up. This result is difficult to interpret due to the unavailability of an MID for SF-12 in the target population. It is possible that our study was simply underpowered to detect a clinically relevant difference. In contrast, the proportion of patients free of subjective dyspnoea clearly decreased in the TERECHO group between a peak at post-treatment and follow-up, returning to about the value at 4 weeks. This suggests that effects on perceived dyspnoea could not be sustained. It is further possible that the post-treatment effect in the participants of the TERECHO group is partly due to their need to reduce cognitive dissonance,90 that is, after the completion of a 6-week exercise programme participants felt the cognitive need to change the perception of dyspnoea even if it had not objectively improved.

The TERECHO programme is targeted at improving physical fitness including physical aspects of subjective HRQOL and should be applied in populations with moderate deficits. Clearly, other patients who have been hospitalised with COVID-19 will be in need of more comprehensive and interdisciplinary programmes. The programme can also not replace early rehabilitation delivered during acute treatment.91 Effects of the programme on pulmonary function seem largely absent while those on mental well-being remain unclear. Components better targeting these outcomes could be added in future evaluations of similar programmes. The TERECHO programme appears to be safe but more mild adverse term effects were found for self-reported dyspnoea and MVV. Effects of the intervention on pulmonary function are otherwise unlikely and effects on mental aspects of quality of life are small at best.

CONCLUSIONS

The TERECHO programme was superior over no rehabilitation with regard to functional exercise capacity, LMS and physical HRQOL. Only short-term effects were found for self-reported dyspnoea and MVV. Effects of the intervention on pulmonary function are otherwise unlikely and effects on mental aspects of quality of life are small at best.

Author affiliations

1Center for Rehabilitation Medicine, Jiangsu Province Hospital/Nanjing Medical University First Affiliated Hospital, Nanjing, Jiangsu, People's Republic of China
2School of Rehabilitation Medicine, Nanjing Medical University, Nanjing, Jiangsu, People's Republic of China
3Department of Rehabilitation Medicine, Hubei Province Hospital of Integrated Chinese and Western Medicine, Wuhan, Hubei, People's Republic of China
4Department of Rehabilitation Medicine, Huangshi Traditional Chinese Medicine Hospital, Huangshi, Hubei, People’s Republic of China
5Institute for Disaster Management and Reconstruction, Sichuan University, Chengdu, Sichuan, People’s Republic of China
6Swiss Paraplegic Research, Nottwil, Lucerne, Switzerland
7Department of Health Sciences and Medicine, University of Lucerne, Lucerne, Switzerland
8XD Group Hospital, Xi’an, Shaanxi, People’s Republic of China

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JL, WX, CZhA, SL and JR designed the study with support from ZY, JW, YC and CZhA; JL led the overall investigation and WX, CZhA, SL and YZ led the investigations at the respective centres; JL acquired the primary funding; WX, CZhA, SL, JW, YZ, YC, CZhA, XF and WC acquired and curated the data with JR responsible for data checks and preparing the data for statistical analysis; JR planned and performed the statistical analysis with support from SL and JW; JR created all figures and tables for this article with support from JW; JR wrote the original draft with support from SL and JW; all other authors revised the draft for critical content; JL and JR supervised the study. All authors have read and agreed with the submitted version of the manuscript. JL and JR act as guarantors for this work and accept full responsibility for the work and the conduct of the study, they had access to the data and controlled the decision to publish.

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Disclaimer

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Competing interests

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Patient consent for publication

No required.

Ethics approval

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Anonymised patient-level data on which the analysis, results and conclusions reported in this paper are based are available as Dryad dataset. Reinhardt, Jan D (2021). Telerehabilitation program for COVID-19 survivors (TERECO) - randomized controlled trial, Dryad, Dataset, https://doi.org/10.5061/dryad.5zw3z27r

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ORCID iD

Jan D Reinhardt http://orcid.org/0000-0001-7250-6906

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