Effect of 12-week pulmonary rehabilitation on cognitive function in patients with stable chronic obstructive pulmonary disease: study protocol for a single-center randomised controlled trial

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ABSTRACT

Introduction Cognitive impairment, an important complication in patients with chronic obstructive pulmonary disease (COPD), seriously affects self-management of the disease and quality of life (QoL). As an exercise-based intervention programme, pulmonary rehabilitation (PR)—especially aerobic exercise (mainly mind–body exercise) and resistance exercise (RE)—has been proposed for its potential effectiveness in improving cognitive function. However, there is still a lack of strong evidence for PR’s effectiveness. In this study, we expect to clarify the effects of pulmonary-based Qigong exercise and elastic band-based RE on cognitive function in patients with COPD and to fill in the relevant evidence blanks.

Methods and analysis This study is a single-centre randomised controlled trial with assessor and data analyst blinding. We will recruit 108 participants with stable COPD starting on 23 December 2019, and randomly allocate them into the pulmonary-based Qigong exercise group, elastic band-based RE group, pulmonary-based Qigong exercise and elastic band-based RE combined group, or control group at a 1:1:1:1 ratio. Participants in intervention groups will perform 30 min of exercise two times per day, 5 days a week, for 12 weeks. The primary outcome will be the global cognitive function as assessed by the Montreal Cognitive Assessment and auditory event-related potential P300. Secondary outcomes will include the specific cognitive domains—attention, memory, executive function, verbal fluency and mental-processing speed, psychological functions and QoL. Exploratory outcomes will include grey matter volume and levels of inflammatory mediators. Outcomes will be measured before and after the interventions.

Ethics and dissemination Ethics approval has been granted by the Ethics Committee of Yue-Yang Integrative Medicine Hospital, an affiliate of Shanghai University of Traditional Chinese Medicine, Shanghai, China (Grant No. 2019-141). Written informed consent will be obtained from each participant before any procedures are performed. The findings will be published in peer-reviewed journals and presented at academic conferences.

Strengths and limitations of this study

► Compare and contrast the feasibility and outcomes following three exercise interventions using a robust randomised controlled trial design and methodology, and include a multidimensional assessment of cognitive function in patients with chronic obstructive pulmonary disease covering global cognitive function (GCF), attention, memory, executive function, verbal fluency and mental-processing speed, and grey matter volume.

► In this study, the main interventions do not require costly specific instruments or much space and can therefore be used for patients who exercise in the community or at home.

► This study combines the Montreal Cognitive Assessment with auditory event-related potential P300 to objectively and precisely assess GCF.

► Assessment results might be affected by subjective bias from the participants in this study because it would be impossible to blind them to exercise as an intervention.

► The effective resistance of the elastic bands cannot be accurately measured during the exercise, and we will strictly control and calibrate the actions of the participants through face-to-face teaching and video learning to achieve the purpose of effective resistance.

Trial registration number ChiCTR1900026869; pre-results.

INTRODUCTION

Chronic obstructive pulmonary disease (COPD) is a systemic disease that impairs physiology, psychology and cognition. Multiple studies with different sample sizes and cognitive-function assessment tools have determined that the prevalence of
cognitive impairment (CI) in patients with COPD ranges from 17% to 56.7%, which is significantly higher than that in healthy controls (12.2%–12.7%). CI in patients with COPD is mainly manifested in attention function, memory function, executive function, verbal fluency and mental-processing speed, which affect physical performance, psychological functions and quality of life (QoL). However, relevant research on the treatment of CI in patients with COPD is rare.

Pulmonary rehabilitation (PR), including but not limited to exercise, health education and behaviour change training, is a comprehensive, multidisciplinary, non-pharmacological treatment based on medical assessment results of patients with COPD. Aerobic exercise (AE) or resistance exercise (RE) at moderate intensity, combined with breathing, is conducive to COPD recovery and plays a positive role in improving cognitive function in patients with COPD. In AE, mind–body exercises such as traditional Chinese exercises (TCEs) can significantly improve the QoL of patients with COPD compared with walking training. Our previous studies also found that TCEs significantly improved lung function, exercise capacity and QoL in patients with COPD. Moreover, a study that applied 6-month Baduanjin for patients with COPD found significant improvements in global cognitive function (GCF; measured by Mini-Mental State Examination (MMSE), as well as by P300), psychological functions (measured by the Hamilton Anxiety Scale (HAMA), and the Hamilton Depression Scale (HAMD)), and QoL. These findings suggest that TCEs have a positive effect on GCF in patients with COPD. TCEs, including Liuzijue, Baduanjin, Wuqinxi and Yijinjing, were organized by the General Administration of Sport of China on the basis of ancient Chinese medicine books in 2003 and are popularised nationwide. They have different characteristics, but only some of them are focused on improving lung function. Therefore, this study rearranged TCEs in order to improve cognitive function in patients with COPD from the perspective of improving lung function; we refer to PR, including both pulmonary-based and Qigong exercise. The movements include the ‘HU’ and ‘SI’ exercises in Liuzijue; ‘pushing up the sky to regulate the Triple Warmer (meridian)’ and ‘drawing a bow to shoot a vulture’ in Baduanjin; ‘the crane actions, including the crane spreading its wings and the crane flying’ in Wuqinxi; and ‘cross-armed iron staff’ in Yijinjing.

RE, as a load-training method targeted at specific muscles, can improve selective attention, associative memory and functional plasticity of the brain in the elderly. Elastic band-based RE can enhance GCF and delay decline in cognitive function in the elderly. The above mentioned results imply that the advantages of RE in CI for patients with COPD are worth exploring. Also, a previous study has shown that a 4-week AE and RE combination regimen significantly improved long-term memory, apraxia and logical-reasoning ability compared with AE alone. Another study compared the effects of AE and RE combination regimen with those of RE alone on executive function and attention in men and found similar results. However, a third study found no difference between AE, RE, and AE combined RE after 12 weeks in improving GCF (as measured by the Montreal Cognitive Assessment (MoCA)) in the elderly. Therefore, it is necessary to explore the advantages of AE and RE combination regimen on cognitive function in patients with COPD.

The main purpose of this study is to explore the effects of 12-week regimens of pulmonary-based Qigong exercise, elastic band-based RE, and the combination of both exercise interventions on GCF, specific cognitive domains (attention, memory, executive function, verbal fluency and mental-processing speed), psychological functions, and QoL in patients with stable COPD. Its secondary purpose is to explore the possible mechanism by which exercise intervention improves cognitive function in COPD, based on brain grey matter volume (GMV) and levels of blood inflammatory mediators. We hypothesised that all three PR regimens are positive significantly, and that combined exercise emerges additional benefits in patients with COPD compared with single exercise methods.

**METHODS**

**Design**

This is a single-center randomised controlled trial with assessor and data analyst blinding. Participants will be enrolled at the Respiratory Medicine Department of Yue-Yang Integrative Medicine Hospital, an affiliate of Shanghai University of Traditional Chinese Medicine. Participants will be randomly assigned to the pulmonary-based Qigong exercise group (QG), elastic band-based RE group (RG), pulmonary-based Qigong exercise and elastic band-based RE combined group (QRG), or control group at a 1:1:1:1 ratio. All participants will receive usual medical care and health education three times for 12 weeks; those in the intervention groups will receive pulmonary-based Qigong exercise, elastic band-based RE or a combination of both exercise. The primary outcome will be GCF as assessed by MoCA and auditory event-related potential (AERP) P300. Secondary outcomes will include the specific cognitive domains as assessed separately by the attention subtest (AT) of the MoCA, Rivermead Behavioural Memory Test-II (RBMT-II), Trail Making Tests Part A and B (TMT-A, TMT-B), Verbal Fluency Test (VFT) and Digit Symbol-Coding Test (DSC) in Wechsler Adult Intelligence Scale (WAIS); psychological functions as assessed by HAMA and HAMD; QoL as assessed by St. George’s Respiratory Questionnaire (SGRQ). Exploratory outcomes will include GMV and levels of inflammatory mediators. Outcomes will be measured within 1 week before and after the interventions. A flow diagram of the study design is shown in figure 1. The study protocol strictly follows the Standard Protocol Items: Recommendations for Interventional
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Trials (SPIRIT) checklist (online supplemental file 1) and SPIRIT figure (online supplemental file 2).

Patient and public involvement

No patients and/or public has been involved in the design, or conduct, or reporting, or dissemination plans of our research.

Sample size

This is a superiority trial. The sample size is calculated based on the MoCA score using the PASS V.11.0 software. The means and SDs of MoCA in the control and AE and RE combined group were 21.20±2.11 and 24.60±2.26, respectively. Setting alpha=0.025 (one-sided), power=0.90, superiority margin=1.13, we calculated the minimum sample size of 21 participants per group. Furthermore, we conducted a power analysis to estimate the sample size needed to perform four group comparisons provided that an α is set as 0.008. The results showed that recruitment of 27 participants per group is a reasonable sample size for this study.

Participants

Inclusion criteria

Participants will be included if they (1) are diagnosed with COPD according to the 2019 Global Initiative for Chronic Obstructive Lung Disease criteria; (2) are clinically stable in the 4 weeks before randomisation (confirmed by absence of acute exacerbation and related drug changes); (3) are 40–80 years of age and (4) are willing to give written informed consent.

Exclusion criteria

Exclusion criteria are as follows: (1) coexistence with other respiratory diseases, including asthma, tuberculosis, bronchiectasis and lung cancer; (2) have been suffering from a severe medical condition that might affect exercise capacity, such as the heart, brain, blood vessels, liver, kidneys, endocrine system, nervous system or musculoskeletal system that might affect exercise capacity; (3) participation in regular exercise training (at least two times per week) in the past 6 months; (4) have been suffering from brain tumour, brain trauma or other diseases that could cause cognitive impairment or (5) presence of a severe medical condition that makes the individual unable to follow instructions properly.

Recruitment and screening

The source of recruitment is the Respiratory Medicine Department of Yue-Yang Integrative Medicine Hospital, where a large number of patients guarantee that enough participants can be recruited. The recruitment will begin on 23 December 2019. The recruitment procedures are as follows: First, the primary researchers will introduce the purpose and the inclusion and exclusion criteria of this study to the recruiting researchers. Then, the recruiting researchers will introduce the purpose, content, and possible risks and benefits of this study to the subjects who have been diagnosed with COPD. If the subjects are interested in participating in the study, the recruiting researchers will talk with subjects in detail to confirm whether they meet the inclusion and exclusion criteria. Finally, the eligible potential subjects or their families will provide signed informed consent to protect their privacy.

Randomisation and blinding

To ensure that the risk of bias remains low, participants will be randomly allocated to one of four groups by an independent person not participating in any other aspect of this study, according to a random number table generated by SPSS V.24.0. Random numbers will be placed in opaque envelopes to achieve allocation concealment, and only the primary researchers will know the randomly allocated sequence. In addition, the assessor and data analyst will be blinded, and they will be replaced once they know the allocation. The new assessor will continue to carry out the assessment, and the new data analyst will re-analyse...
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all data. Meanwhile, we will conduct standard teaching to the assessor through unified learning and practical training within 1 week before the initial and the final assessment, and participants will be strictly required to not discuss interventions with the assessor.

Intervention
All participants will receive usual medical care and monthly health education sessions. Each health education session will last for 60 min and cover the concept, risk factors and self-management of COPD. The control group will be told to maintain their extant daily habits, while the intervention group will be instructed by professional physiotherapists to master specific exercises 1 week before the formal intervention. Interventions will be conducted two times per day (morning and afternoon) for 30 min, 5 days a week. During 1 of these 5 days, participants will gather at the hospital to receive intervention and get feedback; they will exercise at home for the remaining 4 days. The training load will be monitored using Borg CR10 scale at a level of 4–6. If one always cannot perform the specific exercise at the right intensity, then just perform at best-achieved intensity. And the researchers will record it.

Pulmonary-based Qigong exercise group (QG)
Participants in the QG will receive a 12-week intervention of pulmonary-based Qigong exercise (Figure 2). If the Borg reported is below 4 or over 6, participants will be instructed to increase/decrease their intensity, which includes changing the range of motion, speed and breathing method.

Elastic band-based RE group (RG)
Participants in the RG will receive a 12-week intervention of elastic band-based RE. We will use TheraBand elastic bands (The Hygenic Corporation, Akron, Ohio, USA) to provide additional resistance for progressive-RE. Different colours indicate different resistance: brown < yellow < red < green < blue < black < silver < gold. The resistance difference between two adjacent colours is 20%–30%, which will allow a suitable level of resistance for each participant.

In the elastic band-based RE regimen, four upper-limb resistance movements performed by proximal muscles have been selected to improve breathing, and four lower-limb resistance movements to improve walking (Figure 3). Before the formal intervention, one repetition maximum (1RM) that the participant can bear in one exercise will be obtained via the submaximal test (10RM=75% 1RM). The equipment for the 10RM test is the TheraBand elastic band (based on the TheraBand force-elongation table), and the exercises in the test are eight resistance movements in the elastic band-based RE regimen. Referring to similar literature and making appropriate adjustments, exercise intensity will be 30%–40% 1RM in the 1st month, 40%–50% 1RM in the 2nd month and 50%–60% 1RM in the 3rd month. Each session will be composed of 3 sets of 8–12 repetitions of each exercise, with a rest period...
of 1–3 min between sets. If the Borg reported is below 4 or over 6, participants will be instructed to increase/decrease their intensity, which includes changing the colour of the elastic band (including add extra elastic band), repetitions and speed.

Pulmonary-based Qigong exercise and elastic band-based resistance exercise combined group (QRG)

Participants in the QRG will receive a 12-week intervention of pulmonary-based Qigong exercise combined with elastic band-based RE. Total duration of intervention is the same as QG and RG. Participants in this group will be training pulmonary-based exercise for 30 min and elastic band-based RE for 30 min every day.

Outcome measurement

Basic characteristics

Pulmonary function

Specialists will perform the pulmonary-function test using a spirometer (MasterScreen PFT; Jaeger Toennies Erich Jaeger GmbH, Höchberg, Germany) and referring to American Thoracic Society/European Respiratory Society guidelines.32 Each participant will be tested three times, and the final result used in the analysis will be the best value.

Dyspnoea symptoms

The subjective severity of dyspnoea in participants will be measured using the modified Medical Research Council Dyspnoea Scale (mMRC).1 The mMRC is a 5-point scale
(0–4) on which higher scores indicate worse levels of dyspnoea.

Primary outcome
GCF will be measured using MoCA and AERP P300. The MoCA covers 8 aspects of cognitive areas: visual space and execution, naming, memory, attention, language, abstraction, delayed recall and orientation.33 A total score of 30 points and a score of ≥26 points indicate no impairment of cognitive performance. The P300 will be recorded using a 64-channel electroencephalogram signal acquisition recorder (BrainCap MEG; Brain Products, GmbH, Gilching, Germany). A total of 300 stimuli will be presented at 80–85 dB according to the ‘oddball’ paradigm. The ‘oddball’ (novelty/rare) tone will be set at 2 kHz, accounting for 20%, while the standard tone will be set at 1 kHz, accounting for 80%. The bandpass filter will be set in the range of 0.1–50 Hz. Participants will be required to count instances of the ‘oddball’ tone. Brain Vision Analyzer software V.2.2.0 (Brain Products) will be used for the final analysis of the signals collected.34 35

Secondary outcomes
Secondary outcomes include the specific cognitive domains (ie, attention, memory, executive function, verbal fluency and mental-processing speed), psychological functions and QoL.

► Attention function will be assessed using the AT of the MoCA. This test includes three items, with a total score of 7 points. Higher scores indicate better attention.36
► Memory function will be assessed using the RBMT-II that contains 13 items.37 Two parallel versions of the RBMT-II with the same test modes but different materials will be used at the start and end of the intervention. Total score is 24; lower scores indicate greater impairment.
► Executive function will be assessed using the TMT, including the TMT-A and TMT-B. Part A requires participants to connect 25 numbers in order; Part B requires them to alternately connect 25 numbers in squares and circles in order.38 Completion time will be recorded.
► Verbal fluency will be assessed using the VFT, which requires participants to speak the names of animals, vegetables or fruits aloud as often as possible within 60 s.39 The number of the names spoken will be recorded.
► Processing speed will be assessed using the DSC subtest of the WAIS.40 Participants will be required to turn the numbers in the list into corresponding symbols as quickly as possible from left to right within 90 s. Each correct symbol will be scored 1 point; an inverted symbol will be scored 0.5 points. Maximum score is 90 points.
► Psychological functions will be measured using HAMA and HAMD that participants will complete via self-assessments. HAMA will be used to assess anxiety symptoms, which includes 24 items and rated 0–4. HAMD will be used to assess depressive symptoms, which includes 24 items and rated 0–2 or 0–4. The higher the score, the more severe the degree of anxiety and depression.41–43
► QoL will be measured using SGRQ that is a COPD-specific assessment tool.44 The SGRQ contains 50 items, covering symptoms, activities and impacts. The score ranges from 0 to 100, 0 indicating no impairment, and 100 indicating extreme impairment.

Exploratory outcomes
Grey matter volume
Specialists will test GMV using a Magnetom Verio 3.0 Tesla MRI system (Siemens, Munich, Germany). MRI is a non-invasive method of examining brain activity and structure.45 We will use T1 weighted-image 3-dimensional magnetisation-prepared rapid-acquisition gradient-echo sequence to collect T1-weighted structural images.

Inflammatory mediators
The inflammatory mediators including interleukin-6 and 8 (IL-6, IL-8), tumour necrosis factor-α (TNF-α), C reactive protein (CRP) and brain-derived neurotrophic factor (BDNF) will be assayed. Plasma and serum samples will be stored at −80°C as required.

Monitoring of participant compliance
Physical activity and energy expenditure of participants will be assessed using a triaxial accelerometer (ActiGraph GT3X+; Actigraph LLC, Pensacola, Florida, USA) at the start and end of intervention. Participants will be instructed to carry the accelerometer on an elastic belt on the right side of the hip for 7 consecutive days; they will be allowed to remove it during sleeping hours and personal grooming. The threshold of valid data will be set at 10 hours/day. During the test, participants will be required to record the time and reason for removing the accelerometer.

Safety
During the trial, any adverse events will be recorded in a timely manner and reported to the primary researchers, who will then decide whether these adverse events are related to the trial. When a serious adverse event occurs, the Ethics Committee and a clinician from the Department of Respiratory Medicine of Yue-Yang Integrative Medicine Hospital will be informed so that they can provide treatment in a timely manner. For patients who cannot continue training for other reasons, the researchers must record the reason for the drop-out in the case report form, keep in touch with the subjects and complete the assessment as much as possible, and fill out the trial end form.

Monitoring of research quality
Compliance of participants in the control group will be judged based on whether they change drugs due to disease or whether they participated in other regular physical exercises in the preceding 12 weeks. Compliance of participants in the intervention groups will be monitored...
using the exercise recording cards collected during the weekly intensive exercise. Participants whose actual exercise sessions account for >85% of the estimated number of exercise sessions will be judged as having received a valid intervention.

**Data management**
All medical records and assessment data collected from the participants will be securely saved to an encrypted mobile hard drive. Only authorised research assistants will have access to the trial data. Blood samples will be stored and processed in specialised laboratories as required.

**Statistical analysis**
An independent statistician who is not involved in outcome measurement will perform statistical analysis using SPSS V.24.0 software. Intention-to-treat analysis will be used for all randomised participants. Missing data will be analysed according to the likelihood estimation in the mixed effects model. Continuous data will be expressed as mean±SD or median±quartile interval, and enumeration data will be expressed as a relative ratio. Comparisons of baseline differences between groups will be performed using one-way analysis of variance (ANOVA) or the Kruskal-Wallis H test for continuous variables and Χ² for categorical variables. Comparisons within each group (before and after 12 weeks of intervention) will be evaluated using a paired t-test or non-parametric test. Comparisons of endpoint differences between groups will be assessed using a 2×2 ANOVA (group×time) or the Kruskal-Wallis H test. Multiple comparisons between the intervention arms will be performed using SNK-q test or Nemenyi test. P value of <0.05 will be considered statistically significant.

**Ethics and dissemination**
This trial will be conducted in accordance with the Declaration of Helsinki. Approval of this study was granted by the Ethics Committee of Yue-Yang Integrative Medicine Hospital (Grant No. 2019–141). All participants will be fully informed of the research content and the possible disadvantages and benefits of this study, and each will sign an informed consent form before participating in the trial. Study results will be disseminated through peer-reviewed publications and at academic conferences.

**DISCUSSION**
A variety of influencing factors may cause CI in patients with COPD, including age, disease duration, disease severity, hypercapnia, smoking and vascular dysfunction.45–47 Of those, hypoxia caused by pulmonary dysfunction is the main cause of CI in patients with COPD. The lower the partial pressure of oxygen, the higher the risk of CI.48 49 However, the pathological mechanism remains unclear. There is a hypothesis that prolonged hypoxia produces oxygen-free radicals and then releases inflammatory mediators, which affect the synthesis and metabolism of neurotransmitters and cause glial-cell activation; further damage to neurons leads to changes in brain structure and function and eventually to CI (figure 4).50 51 Previous studies have shown that CI in patients with COPD is associated with abnormal GMV. In one such study, Wang et al found that local GMV in patients with COPD was significantly reduced compared with that in the normal population.44 Yin et al found that the left-superior frontal gyrus and the right orbital part of the inferior frontal gyrus are strongly associated with pulmonary function (as measured by forced expiratory volume in one second) and cognitive function (as measured by MoCA).52 Zhang H et al found that the right-inferior triangular frontal gyrus in particular is related to graphic memory.53 However, the pathological mechanism of GMV atrophy in patients with COPD is very complicated and has not been conclusively demonstrated. In addition, inflammatory mediators might play an important role in the development of CI. In healthy people, higher level of the peripheral blood inflammatory mediator CRP significantly increases the risk of CI,54

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**Figure 4** The possible pathological mechanisms of CI in patients with COPD: pulmonary dysfunction is the main influencing factor; other influencing factors include age, disease severity, hypercapnia, smoking, cardiovascular disease and genetic susceptibility. The underlines will be observed before and after the intervention in this study. BDNF, brain-derived neurotrophic factor; CRP, C reactive protein; IL, interleukin-6; PaO₂, pressure of arterial oxygen; TNF-α, tumour necrosis factor-α.
especially mental-processing speed.55 In patients with Alzheimer disease, there is a meaningful correlation between inflammatory mediators and GCF.56,57 Moreover, it has been shown that the plasma of patients with COPD often contains abnormal levels of inflammatory mediators, including IL-6, IL-8, CRP, TNF-α and BDNF.58–60 Therefore, in this study, we seek to observe whether PR can affect GMV and blood inflammatory mediators and to explore the possible mechanism by which exercise affects cognitive function in patients with COPD.

TCEs are moderate-intensity mind–body exercises that coordinate bodily movement, breathing, posture and attention.61 62 Pulmonary-based Qigong exercise has strong lung-enhancing effects and is more suitable for patients with COPD.63–65 Considering that CI in patients with COPD is closely related to lung function impairment, this study intends to observe whether pulmonary-based Qigong exercise can improve cognitive function in such patients from the perspective of enhancing their pulmonary function.

As an important component of PR, RE can effectively improve physical function and QoL in patients with COPD. Conventional RE relies on large and expensive equipment,63 while elastic band-based RE is easily accessible and does not require much space. In addition, elastic bands can provide different levels of resistance individually. Evidence has demonstrated that elastic band-based RE has the same positive effect as conventional RE (weight machines and dumbbells) in interventions for muscle strength and QoL in patients with COPD.64 65 Therefore, this study intends to observe the effect of elastic band-based RE on cognitive function in patients with COPD.

In terms of cognitive-function assessment tools, we will use the MoCA Scale and AERP P300 to assess GCF in patients with COPD. As a comprehensive, time-consuming, easy-to-use assessment tool verified to be both sensitive and specific, the MoCA has a higher diagnostic efficacy for CI than the similar MMSE Scale.66 67 In both sensitive and specific, the MoCA has a higher diagnostic efficacy for CI than the similar MMSE Scale.66 67 In addition, it has been found that P300 latency in patients with COPD is significantly prolonged,68 and significantly differ by disease duration.54 68 Compared with MoCA, AERP P300 results have the clear advantage that are not affected by subjective factors. However, the specificity of AERP P300 in the diagnosis of CI is not very strong.69 Therefore, MoCA combined with AERP P300 can more accurately and objectively assess cognitive function in patients with COPD.

In summary, this article presents the study protocol for a clinical trial to investigate the effects of pulmonary-based Qigong exercise, elastic band-based RE, and a combination of both exercise on GCF, specific cognitive domains, QoL and psychological functions in patients with stable COPD. The study results will clarify the effects of 3 PR regimens on cognitive function in patients with stable COPD and further explore possible mechanisms by which PR intervention affects cognitive function in these patients.

Trial status
This trial is currently in the recruitment phase.

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Contributors HD and PL are joint first authors. HD, PL, ZW, HC and TW designed the study. HD drafted and organised most of the manuscript. PL drafted part of the manuscript, guided the key revision of the manuscript and participated in trial registration and ethical review. ZW contributed to the participants’ recruitment. HC and TW provided design ideas for the study, collected relevant data and made substantial contributions to the production of pictures in the manuscript. XL and WW contributed to the interpretation of the results and critical revision of the manuscript for important intellectual content and approved the final version of the manuscript. All authors have read and approved the final manuscript.

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