Efficacy of cognitive behavioural therapy-based smartphone app for smoking cessation in China: a study protocol of a randomised controlled trial

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ABSTRACT

Introduction  A number of smartphone apps have been proved to be effective in assisting smoking cessation in many countries, but little is known about the effectiveness of smartphone apps that assist in smoking cessation in China. This study aims to evaluate the efficacy of cognitive behavioural therapy (CBT)-based smartphone smoking cessation smartphone app for treatment-seeking smokers in China.

Methods and analysis  In this remote control, randomised controlled trial, the participants will be randomised and allocated in 1:1 ratio to smoking cessation app intervention group or to control group. Participants will receive a 12-week intervention (treatment group will receive CBT-based cigarette smoking cessation smartphone app intervention and control group will receive text messaging-based control intervention) with follow-up at week 26. The primary outcome includes biologically verified continuous smoking abstinence at week 26 after the quit date. The main secondary outcomes include 7-hour point prevalence smoking abstinence at weeks 1, 2, 3, 4, 8, 12, 16, 20 and 26, self-reported continuous smoking abstinence at weeks 4, 8, 12, 16, 20 and 26 after the quit date. About 1200 smokers with willingness to make a quit attempt within a month from January 2021 to August 2021 or until the recruitment process is complete will be recruited. The final 26-week follow-up will be completed in August 2022. The trial results will be available by the end of 2022. All analyses will be done on an intention-to-treat basis. Based on the previous studies, we hypothesised that this CBT-based app will be an effective, feasible and accessible smoking cessation app in China.

Ethics and dissemination  Ethical approval was obtained from the ethics committee of Sir Run Run Shaw Hospital, an affiliate of Zhejiang University, Medical College (number: 20200129-33). Study results will be disseminated via research conference presentations and peer-reviewed publications.

Trial registration number  NCT04421170.

INTRODUCTION

Smoking is the leading cause of preventable morbidity and mortality throughout the world, especially in China with approximately 316 million smokers, and this accounts for nearly one-third of the world’s total smokers.1–3 In October 2016, China has announced a new national strategy, that is, Healthy China 2030, which involved decrease in cigarette smoking rate to 20%.4 The WHO Framework Convention on Tobacco Control has taken effective measures to control tobacco use and provide adequate treatment for tobacco dependence. However, smoking cessation rates remain low in China.5 Insufficient smoking cessation services are one of the most important contributing factors for the low cessation rates that have been reported in China.6 Our previous randomised controlled trial (RCT) of text-messaging-based smoking cessation intervention (‘Happy Quit’) has shown an increased biochemically verified continuous smoking abstinence at week 24 from around

Strengths and limitations of this study

▶ The smoking cessation app is the first to be developed based on the theoretical frameworks guided (cognitive behaviour theory), evidence-based randomised controlled trial by adhering to the guideline recommendations in mainland China.

▶ If this app programme is proved to be effective, then it will have a huge potential use in China, where in there are more than 300 million smokers, at population-level, and the public health impact could be enormous.

▶ Abstinence will be mainly measured via self-report through app questionnaires, which is less rigorous than a biochemically verified abstinance. Thus, this study will provide biological verification for those individuals who have self-reported continuous smoking abstinence at 6 months and chose this as the primary outcome in this study.

▶ Participants in this study will be more likely motivated than other smokers, and they can search for other smoking cessation apps or any other available smoking cessation services during the trial.

▶ Participants can drop out from the trial at any time point, as this smoking cessation app programme provides only online services.
2% to more than 6%. The provision of more comprehensive interventions, such as widely accessible online treatment strategy for smoking cessation, might further increase the quitting rates.

So, finding more effective strategies for treatment-seeking cigarette smokers to quit smoking has become a priority due to high smoking rates in China. With increasing popularity of smartphone usage in Chinese general population, mobile devices have become the most common ones in healthcare settings for promoting health, and this led to a rapid growth in the development of medical software applications (hereafter referred to as ‘apps’) for these platforms. These smartphone apps provide support at a very low unit cost for those individuals who wish to quit smoking. It has become a promising way for smokers, especially the new generation smokers who might be more open to receive cessation support via digital technologies.

Currently, there are numerous apps available that assist in smoking cessation, but most of these did not adhere to the clinical practice guidelines for treating tobacco use and dependence and did not demonstrate clinical efficacy. These might be because of the ‘low-certainty evidence of the benefit of smartphone apps interventions’ according to a Cochrane Systematic Review. During the past decade, a variety of smartphone apps (approximately 70 apps) have been developed for smoking cessation in China. However, few of these have adhered to the China Clinical Smoking Cessation Guidelines (issued in 2007 and updated in 2015), and these are outlined directly from the US Clinical Practice Guideline for Treating Tobacco Use and Dependence. These available Chinese apps neither referred the user to a recommended treatment nor connected the users to anything outside the app, like quitline counselling or approved medications. However, the content of these Chinese apps is primarily limited and more than 80% of apps were downloaded for less than 10000 times.

By integrating with the guidelines and other evidence-based practices in a better manner, several studies have indicated that smoking cessation apps are effective ways for reducing smoking prevalence, especially among young adults. As China has huge cigarette smoking population with very limited evidence-based smoking cessation interventions, development of an evidence-based app that adheres to the guidelines acts as a powerful tool for smoking cessation. Furthermore, according to a survey conducted, a majority of healthcare providers have agreed that mHealth apps hold promising features for individuals to quit smoking and would be recommended to their clients, if the programme was empirically validated.

SMOKING BEHAVIOUR IS AN UNHEALTHY BEHAVIOUR THAT CAN BE CHANGED, BUT CHANGING THE BEHAVIOUR OF PEOPLE TO REDUCE SMOKING IS NOT AN EASY THING. PERFORMANCE, STRESS AND BODY WEIGHT ARE THREE MAJOR FACTORS THAT INFLUENCE THE SMOKING BEHAVIOUR OF THE PEOPLE. COGNITIVE BEHAVIOURAL THERAPY (CBT) IS ONE OF THE MOST COMMON AND THE BEST-STUDIED FORMS OF PSYCHOTHERAPY THAT INVOLVES CHANGING MALADAPTIVE THINKING PATTERNS AND THE NEGATIVE BEHAVIOURS ASSOCIATED WITH THEM. CBT IS AN EFFECTIVE WAY TO TREAT CANNABIS AND NICOTINE ADDICTION BUT IS LESS EFFECTIVE IN TREATING OPIOID AND ALCOHOL ADDICTION. CBT IS USED AS A PSYCHOLOGICAL INTERVENTION FOR PEOPLE WHO WANT TO QUIT SMOKING BY CHANGING AND RESTRUCTURING THEIR THOUGHT PROCESSES AND ITS ASSOCIATED BEHAVIOURS, WHICH IN TURN ASSISTS IN EFFECTIVE SMOKING CESATION AND PREVENTS RELAPSE. THE TRAINING OF COGNITIVE BEHAVIOURAL SKILLS ASSISTS SMOKER TO CHANGE THEIR SMOKING BEHAVIOUR INTO NO-SMOKING BEHAVIOURS AND RESTRUCTURING THEIR THOUGHT PROCESSES. IT INVOLVES NEW LEARNING BEHAVIOURS TO EFFECTIVELY QUIT SMOKING, MAINTAIN ABSTINENCE, REDUCE WEIGHT CONCERNS AND COPE WITH DEPRESSION.

This RCT evaluates the efficacy of smoking cessation smartphone app, which is a Mandarin smoking cessation app that integrates smoking cessation support and social skills training into cognitive behavioural principles and behaviour change techniques, in China. Based on the efficacy of previous smoking cessation app studies in other countries and mobile phone-based text message interventions (such as ‘Happy Quit’, based on the CBT) for smoking cessation in China, this CBT-based app is hypothesised to be an effective, feasible and accessible smoking cessation app in China. Also compared with the control app group, the rate of biologically verified continuous smoking abstinence at week 26 after the quitting date will be substantially higher in the CBT-based cigarette smoking cessation app group.

METHODS AND DATA ANALYSIS
Participants and public involvement
Neither the participants nor the public will be involved in the design, recruitment or conduction of this study.

Study design and participants
This internet-based RCT will be conducted in China. About 1200 smokers (a smoker is defined as a person who smoked more than 100 cigarettes in their lifetime, and currently smoke 5 or more cigarettes a day) who are willing to make a quit attempt within a month will be randomly allocated using a randomisation method of Interactive Web Response System (IWR). The smokers based on balance for age (25–34 vs ≥34 years) and cigarettes per day (3–20 vs >20 cigarettes per day) will be randomly allocated to smoking cessation app that comprises of motivational messages and cognitive and/or behavioural change support or to the control group that receives only text messages of thanking them for being part in this trial and reminding them till their free month at the end of follow-up. Only the participants in the intervention group will receive 12-week intervention programme. These participants will be recruited from January 2021 to August 2021 or until the recruitment process is completed. The final 26-week follow-up period will be completed by August 2022. All trial results will be
Inclusion criteria:
1. Cigarette smokers (smoked more than 100 cigarettes in their lifetime, and currently smoke 5 or more cigarettes a day).
2. 25 years of age or older.
3. Being able to read and write in Chinese.
4. Owning a smartphone*.
5. Have experience of using apps.
6. Expressing an interest in quitting smoking within the next month.
7. Willingness to use only the study app during the course of the study.
8. Willing to provide informed consent to participate in the study.

Exclusion criteria:
1. Non-smokers.
2. Only use electronic cigarettes.
3. Smokers without attempts to quit smoking.
4. Participants with severe mental illness†.
5. Participants who had already started their quit attempt or using any smoking cessation treatment at the time of registration.
6. Unable to use smartphone and apps.
7. Unable to read and write in Chinese.
*Smartphone operating system iOS V.9.0 or above; Android V.5.0 or above. Only subjects with iOS smartphone will be recruited during pilot phase.
†Any individuals with psychiatric symptoms that are debilitating that their ability to participate in this trial is severely impaired. These illnesses include schizophrenia, schizoaffective disorder, bipolar disorder and major depression.

available by the end of the year 2022. The overview of the eligibility criteria of the participants is described in box 1.

Sample size and power calculation
Sample size calculations will be done based on the differences in the objective measures of the primary outcome event, that is, biologically verified continuous smoking abstinence at week 26 between intervention and control groups. Based on our previous study of ‘Happy Quit’, a ratio of 1:1 will be assumed for intervention to control participants, with an alpha of 0.05, power of 80% and an effect size equal to 7% in the intervention versus 2% in the control condition, the required sample size is 486 per group with a total of 972 participants by using a two-tailed t-test on proportions. Considering a 20% lost to follow-up in this study, the final required sample size is 600 per group with a total of 1200 participants in both the groups. We planned to assess a preliminary result on 200 participants in a pilot study with 1:1 ratio (100 participants in the intervention group and 100 participants in the control group).

Recruitment
Online advertising will be used to recruit potential participants. This service will be advertised online (such as Weibo or other websites, WeChat, QQ et al) and offline (such as universities, companies, communities, hospitals). Potential participants will register by sending text messages or making a call. The research assistants will then contact the respondents to assess their eligibility and explain the study to each participant and inform them that they will be allocated to either a control group or an app programme. Baseline data and outcome data will be collected by Electronic Patient Reported Outcomes (ePRO) software (http://www.c-path.org/programs/pro). Informed consent will be electronically obtained from each participant using an e-consent system before collecting the baseline data. Participants enrolled in this study can be withdrawn at any point of time. The participants will be asked to provide a mobile phone number for baseline assessment and follow-up as well as for receiving mobile phone-based payment as reward.

Randomisation and group allocation
The eligible participants who provide their consent to participate in the trial and have completed their baseline assessment will be randomly assigned to either intervention or control group. Randomisation (by IWRS) will be based on 1:1 ratio at individual level by balancing sex, age, level of educational, previous quit attempts and smoked cigarettes per day. The study investigators will be blinded to the participants’ treatment allocation until all data have been collected. The investigators assessing the outcomes will be blinded to the participants’ allocated groups until all data are analysed.

Development of CBT-based cigarette smoking cessation smartphone app
This CBT-based cigarette smoking cessation smartphone app will be developed by Johnson & Johnson pharmaceutical company, supported by Neusoft Corporation and supervised by the Department of Psychiatry, Sir Run Run Shaw Hospital, School of Medicine, Zhejiang University. The content of the intervention will be developed based on the principles generated by the CBT and the clinical experience, scientific or empirical evidence of smoking cessation.

Eligible participants will download the app through their smartphones and activate the app by entering the code. In order to provide tailored information, such as motivational message-sending recommender, to each participant based on their personal information, their basic and smoking-related information including age, gender, years of smoking, number of cigarettes per day and motivation and self-confidence regarding smoking cessation will be entered into the app.

Theory
This CBT-based app programme integrates cognitive behavioural principles and behaviour change techniques into individualised smoking cessation support and social skill training. The aim of this app programme involves reconversion of smoking-related thoughts, knowledge and behaviour related to smoking, and it correspondingly changes smoking behaviour into stop smoking behaviour of cigarette smokers. For example, CBT technical will be used to address the smokers’ concerns with regards to smoking/nicotine enhancement in the aspect
of performance and reduced stress or concerns of weight-gain, rather than focusing on preventing the weight gain itself.

**App features**
The design of the app will primarily focus on the ease of use of features to promote engagement and evidence-based smoking cessation practice to improve the abstinence rates of cigarette smokers. A cigarette smoking end-user team will be invited to verify the ease of use and engagement. The overview of the app features that could enhance the ease of use is shown in table 1 and the app features that could promote engagement are shown in table 2.

**Summary of main features**
This CBT-based app involves a scientific-based approach with evidence-based techniques to help the participants quit smoking for good. The summary of the main features of the app are as follows:

1. CBT-based techniques for smoking cessation.
2. Create a personalised plan to help participants quit smoking for good.
3. Choose an approach that works the best for participants: either going cold turkey or slowly decreasing their nicotine intake.
4. Track the participants’ cigarette consumption and their cravings.
5. Track money saving.
6. The app is built-in family and friends’ support.
7. Relaxation techniques to reduce stress.
8. Multiple ways to deal with craving to smoke.
9. Motivational tips.
10. Health achievements (from their blood circulation and oxygen levels to their senses of taste and smell).
11. Emergency SOS to keep participants smoke-free for good. Multiple options will be provided to deal with them.

| Feature | Brief description |
|---------|------------------|
| Navigation | Ease of navigation; users will directly navigate the menu choices |
| Displays | Simplified pleasing displays with light green colour |
| Aesthetics | Keep main pages simple and visually appealing and encourage and make it easy to use |
| Minimum text | Keep text as brief as possible |
| Text formatting | Use simple and easy to understand sentence, use plenty of headings, keep paragraphs short, use bulleted lists and highlight key terms |
| Easy-to-read | Reading level to age 14 (education of middle school level) |
| Language | Clear and consistent |
| Font size | No small text |
| UI button setting | UI button setting design conforms to user habits |

| Feature | Brief description |
|---------|------------------|
| Gamification | Attractive gamified quitting journey with continuity story line and variable plot |
| Personalisation | Make the app as interactive as possible (eg, tailored feedback); promote engagement by customised message push notification based on users’ own smoking behaviour |
| Professionalisation | Make the app looks professional with scientific endorsed content |
| Personification | Establish a ‘rapport’ between the participants and healthcare providers |
| Social interaction and support | Provide a platform for interactions among the participants, their quitting sponsors and healthcare providers. Linking with social networking software that is, WeChat |
| Transparency and realistic expectations | Set up clear expectations concerning how the app will be used |
| Shaping | Keep demands of the participants to a minimum |
| Design for curiosity | Unluck some information each day during preparation; present new information each time the app is accessed |
| Instant feedback | Provide instant feedback (eg, tips for craving, progression statistics) |
| Rewards | Provide users with a rewarding experience when they visit the app |
| Positive recognition | Record quitting time and provide recognition and encouragement by gamified achievement system, visualise health and finance improvement |
| Visual cues and dashboards | Use as many as possible images to convey information |
| Structure sections | Break complex tasks into small steps |
| Session time | Keep login sessions brief (each session takes less than 1 min of the users’ time) |
Evidence-based smoking cessation context in app

The content in app involves CBT targets of both smoking-related cognitive and behavioural problems by fully integrating cognitive and behavioural strategies. The content will be guided by the cognitive behavioural theoretical frameworks and is adapted from the text messaging-based smoking cessation intervention\(^1\) and other evidence-based smoking cessation interventions\(^2\) as well as information from a survey conducted on design and development of smoking cessation apps.\(^1\) Based on this survey, the following information will be obtained from the smokers: sharing the process of smoking cessation with family members and friends (67.7%), help them track their progress (66.8%) and help them with their withdrawal symptoms or side effects of medications (63.0%).\(^1\)

The content contains two major parts:

1. A questionnaire including the data about demographics (gender, age, height and weight for calculating body mass index (BMI), levels of education), smoking behaviour scale of Fagerström Test for Cigarette Dependence (FTCD), previous use of smoking cessation apps and general use of health-related apps.

2. Theoretical framework guide, wherein it includes the evidence-based information with regards to smoking cessation. The app programme content will be based on cognitive behavioural intervention techniques and smoking cessation guidelines that focus on education, cognitive restructuring, self-monitoring and practical coping strategies to increase the participants’ self-efficacy for successful smoking and eventually increasing the abstinence rates. CBT will be performed based on the thoughts/thinking guide of the cigarette smoker’s behaviour. For example, illogical beliefs like enhanced smoking performance (cognitive process) leads to dysfunctional behaviour of continued cigarette smoking (behavioural action). To acquire the capability of quitting smoking, individuals need to recognise, understand and change their smoking behaviour patterns of monitoring their smoking patterns, identify high-risk situations, adapt the coping responses to life stressors and risky situations. The examples of types of information included in different phases of cognitive behavioural smoking cessation app programme are shown in online supplemental table 1. The information in smoking cessation app will adhere to the recommended smoking cessation practices that are presented in table 3.

### Intervention

First, the research investigators will assess each of the eligibility of each participant. At baseline assessment, all eligible participants will be invited to complete a questionnaire that contains items on demographic and smoking-related characteristics such as previous quit attempts, current smoking behaviour, motivation and determination to quit, smoking dependence by ePRO software. After completing the baseline questionnaire,
eligible participants will be automatically randomised to either intervention app or control app. The questionnaire will take about 5–10 min to complete. Randomisation will be done by Electronic Data Capture (EDC) software. The participants from both groups will be motivated to set a quit date at 1 week. The participants will receive a programme (prequit: 1 week and postquit: 12 weeks) of either personalised information (intervention group) by smartphone app or control information (control group) by mobile text messaging (short message service (SMS)) from the day after completing the questionnaire, and then will be followed up until week 26 postquit date. Both groups will have no restriction to seek other cessation methods.

Control group
For participants in the control group, SMS will be sent as are minder to fill out the self-report questionnaire via ePro system until week 26 after the quit date. After this period, the SMS will acknowledge and recommend participants to receive intervention by app for free if they want. The participants who are allocated to the control group will be encouraged to make a quit attempt and select a quit date at 1 week after completing the baseline questionnaire. They will not receive any other smoking cessation information and will receive information of thanking them for being in the study and reminding them of the time until their free month at the end of follow-up. To measure the outcomes between the two groups, continuous smoking abstinence, point prevalence of abstinence, how many cigarettes per day during the last week if they are still smoking will be checked at the end of weeks 1, 2, 3, 4, 8, 12, 16, 20 and 26 points after the quit date by ePRO software. Biochemically verified continuous abstinence will also be checked if they have reported continuous smoking abstinence at week 26 after the quit date.

Intervention group
The participants in the intervention group will receive CBT-based smoking cessation and details about the intervention app as shown in ‘2.5 Development of CBT-based cigarette smoking cessation smartphone app’. This will provide both mandatory information of evidence-based and guideline-based smoking cessation interventions, and optional information on quitting benefits, tips for quitting and so on. The app will provide the participants (1) with several tasks/information each day, (2) free to view any information, (3) automatic notifications at different stages of quitting process, such as daily motivational messages, tips for quitting, (4) participants can also seek for help, (5) encouragement to monitor and record their smoking behaviour or write quitting diaries (eg, craving triggers), (6) to include a quitting benefits tracker that will notify the user about the health benefits and financial benefits gained after quitting and (7) to monitor their daily usage data. The app will be available for participants in the intervention group until week 26 postquit date follow-up. After this period, the app will automatically stop collecting the data but can be used if they wish to. The smoking cessation information will be tailored according to the responses provided by the participants in the smoking questionnaire and include review of their personal smoking behaviour, remind their reasons for quitting, tips to support their quit attempt, encourage to increase their motivation and boost their self-confidence, support them on how to cope with high-risk situations when they are tempted to smoke and remind them to get social support. More information will be provided during the preparation phase and early postquit phase (the first 4 weeks, especially the first week). As the participant’s progress throughout the study, smoking cessation-related information will be gradually reduced until week 12 after quit date, and follow-up messages will be sent at weeks 16, 20 and 26 after the quit date. If the participants have any questions on using the smoking cessation app, they can also seek for help (from Dr Liao, Dr Tang or research assistants, will be documented) at any time by text or WeChat or make a phone call.

Outcomes
1. Primary outcome:
   a. Biologically verified continuous smoking abstinence at week 26.
2. Secondary outcomes:
   a. Seven-day point prevalence of smoking abstinence at weeks 1, 2, 3, 4, 8, 12, 16, 20 and 26 follow-up after the quit date.
   b. Self-reported continuous smoking abstinence at weeks 4, 8, 12, 16, 20 and 26.
   c. Reduction in the number of cigarettes smoked per day.
   d. Rates of participation in and completion of smoking cessation programmes.

Table 3  Continued

| Guideline recommended context/5 As | Yes (✓) | No (X)* |
|-----------------------------------|---------|---------|
| Discussion about pharmaceutical products | X |         |
| Referral to quitline or other professional support | X |         |
| Support for craving and withdrawal symptoms | ✓ |         |
| Arrange follow-ups |         |         |
| Checked-in prior to and after quit attempt | ✓ |         |
| Support for high-risk situations | ✓ |         |
| Support for craving and weight concern | ✓ |         |
| Support for lapse and relapse prevention | ✓ |         |
| If relapsed, encourage user to set a new quit date | ✓ |         |
| If relapsed, offer encouragement that quitting takes practice | ✓ |         |
| Support identity change | ✓ |         |
| Reward and support long-term abstinence | ✓ |         |

5 A’s: Ask, Advise, Assess, Assist, and Arrange
* (X) This information will not be provided until the end of clinical trial.
Programme acceptability will be assessed at week 4 postquit (pilot phase).

**Measures**

1. Biologically verified continuous smoking abstinence: Participants in this study will be considered to be biologically verified continuous smoking abstinence if they report smoking of no more than five cigarettes since the quit date and have a urine cotinine cut-off point of 200 ng/mL. Cotinine urine dipsticks will be mailed to each participant's address, and test will be confirmed by video call and by family members) at week 26 (6 months) after the quit date. This abstinence has been applied by the Society for Research on Nicotine and Tobacco (SRNT) and the 'Russell Standard'.

2. Seven-day point prevalence of smoking abstinence: participants who did not even have a puff of smoke for the last 7 consecutive days, and at weeks 1, 2, 3, 4, 8, 12, 16, 20 and 26.

3. Self-reported continuous smoking abstinence: a self-report of smoking ≤5 cigarettes from the past till weeks 1, 2, 3, 4, 8, 12, 16, 20 and 26.

4. Cigarettes smoked per day: number of cigarettes smoked per day within 7 days or the approximate number of cigarettes smoked within 7 days if not smoked daily.

5. Reduction in the number of cigarettes smoked per day: will be assessed by comparing the number of cigarettes smoked per day at baseline and at week 26.

6. Nicotine dependence: nicotine dependence will be measured by the Chinese version of the FTCD. Nicotine dependence is classified using the FTCD score with the following categories: very low/no dependence (FTCD score: <4), low to medium dependence (FTCD score: 4–6) and high dependence (FTCD score: ≥7).

7. Smoke craving: smoke craving will be measured by visual analogue scale (VAS, 0–10 integers), which includes measurement of craving for a cigarette.

8. Self-efficacy: self-efficacy will be measured by VAS measurements of participants’ readiness and confidence to quit smoking.

9. Motivation to quit: motivation to quit will be assessed by VAS measurements of participants’ motivation to quit smoking.

10. Programme acceptability: programme acceptability will be evaluated by questions for assessing programme acceptability in **Table 4**. A small sample (approximately 100 participants) will be assessed at week 4 after the quit date by ePRO software.

**Table 4 Questions for assessing ‘CBT-based smoking cessation app’ programme acceptability**

| Question | Rating |
|----------|--------|
| 1. Overall rating of the programme | Like very much<br>Like somewhat<br>Neutral<br>Dislike somewhat<br>Very dislike |
| 2. Appraisal of ‘CBT-based smoking cessation app’ programme; likelihood of recommending programme to others | Very likely<br>Somewhat likely<br>Neutral<br>Unlikely<br>Not at all likely |
| 3. The experience of using app: ease-of-use | Strongly agree<br>Agree<br>Neutral<br>Disagree<br>Strongly disagree |
| 4. The experience of using app: promoting engagement | | |
| 5. The programme made it easier to quit smoking | | |
| 6. The programme is not the one I was expected | | |
| 7. I would not have been able to quit without the programme | | |
| 8. I stopped using it by the end of the programme | | |
| 9. Information was easy to understand | | |
| 10. Information was useful for me | | |
| 11. The information mentioned about what I was experiencing and feeling | | |
| 12. I received too much information | | |
| 13. Frequency of using app | Almost never<br>≥1 time a week<br>Everyday |

CBT, cognitive behavioural therapy.

follow-up). The drop out rates at each visit will be at least below 25%.

- Response rates: the percentage of responses for questions with regards to outcomes at each time.

**Procedures**

**Figure 1** shows the trial flow chart. Participants will be assessed at baseline (0 week), at weeks 1, 2, 3, 4, 8, 12, 16, 20, and followed up at week 26 after the quit date. The schedule of enrolment and follow-up assessments are summarised in **Table 5**. The demographic characteristics of gender, age, height and weight (for calculating BMI), educational level (primary, middle school, high school, university and above) will be included. Smoking characteristics of cigarettes per day, quit history, motivation and confidence to quit, quitting smoking gradually or abruptly, nicotine dependence by FTCD and smoking craving by VAS will be included.

Standard abstinence questions will be asked: ‘Have you ever smoked during the past 7 days?’ and ‘Have you ever...
smoked since (the date of start of abstinence period, that is, from week 2)’ A: No, not a puff; B: 1–5 cigarettes; C: More than five cigarettes?’ at weeks 1, 2, 3, 4, 8, 12, 16, 20, and at follow-up week 26 after the quit date. If a participant’s response is A or B at all points, then a biochemical test will be conducted to confirm the individual as an abstinent. If the response is C, questions regarding cigarette consumption will be asked. Questions about weight gain will be asked monthly until the end of the follow-up week 26 after the quit date. Weight change will be recorded monthly until 6 months postquit. At the end of the trial, participants will be asked, ‘Have you used any other smoking cessation methods: such as visit smoking cessation clinics, consulting, smoking cessation medications like nicotine replacement therapy, bupropion, varenicline or any other smoking cessation services or medications’.

Withdrawal from the programme
Participants will be informed to choose to stay in the app or not without giving any reason. Also, an individual who initially agrees to participate in this study will have the right to withdraw from the study at any point of time and have the right to refuse to answer any particular question(s) or participate in a particular set of procedures in this trial. All participants who still stay in the app will be contacted and not required to provide withdrawal consent, and will be followed up irrespective of their smoking status. Based on the intention-to-treat (ITT) principle,31 participants who fail to respond to the follow-up end-point will be retained in the analyses and classified as smokers. However, if participants are unwilling to participate in the study, then they will be excluded from the analyses and the reason(s) for their withdrawal will be recorded if provided.

Data collection
Questionnaire data will be automatically collected by a data acquisition system into the ePRO Software (http://www.c-path.org/programs/pro). A user-specified file will be downloaded from the database, and the data will be subjected to basic processing and recoding. There will be only one interim analysis with the 200 participants initially...
for assessing the programme acceptability. Only investigators can access data. All data will be monitored by the data monitoring committee of the hospital. Personal data will be deidentified.

**Statistical analysis**

On completion of data collection, the data will be cleaned up and prepared for data analysis. The primary and secondary outcome variables and mediating mechanisms will be checked for distribution, outliers and missing patterns and appropriate steps will be taken as and when necessary. All analyses will be conducted based on an ITT. In order to address the non-responses at follow-up, the missing data from lost to follow-up will be expected to be a smoker.

Demographic and smoking-related characteristics will be compared between the groups at baseline using independent $t$-test for continuous variables or $\chi^2$ test for binary variables as appropriate. Logistic regression will be used for between-group comparisons of the primary outcome variable—biologically verified continuous smoking abstinence at week 26 and the secondary outcomes 7-day point prevalence of smoking abstinence (at weeks 1, 2, 3, 4, 8, 12, 16, 20 and 26) and self-reported continuous smoking abstinence (at weeks 4, 8, 12, 16, 20 and 26). For comparisons of secondary continuous outcomes that do not meet the normal distribution assumptions, a non-parametric Mann-Whitney test will be conducted. To account for the non-responses at follow-up, four multiple imputation models will be constructed for the non-responses during the follow-up at 7 days, week 4, week 12 and week 26 of continuous abstinence. The covariates included in the models are age, gender, education level, nicotine dependence and intervention group. Ten imputed data sets will be generated based on Rubin’s formula for relative efficiency to produce about 99% efficiency. In addition, a complete case analysis will be performed, in which any participant with missing information on any outcome will be excluded. Finally, a subgroup analysis will be conducted for key demographics, smoking and cessation characteristics, social support and use of intervention variables to assess homogeneity of the treatment effects using logistic regression. The effect measures include relative risk and 95% CIs. Further, a two-sided $p$ value of $<0.05$ will be considered as statistically significant difference. The completed data will be analysed and blinded to the intervention assignment by a trial statistician using the R software (R Foundation for Statistical Computing, Vienna, Austria. https://www.r-project.org/) and the IBM SPSS Software (IBM, IBM SPSS Statistics for Windows, https://www.ibm.com/analytics/spss-statistics-software). Also, a sensitivity analysis will be conducted with the assumption that all participants with missing outcome data had not changed the smoking status since the previous follow-up (ie, ITT).

**Safety**

Throughout the ‘CBT-based smoking cessation app’ programme, the participants from both groups will be encouraged to communicate with us (Dr Liao, Dr Tang or research assistants) if they experience any smoking cessation relative health issues. The contact information (available at any time) will be provided to each participant at baseline. Most of the nicotine withdrawal symptoms are only temporary and subside within several weeks. If participants experience severe physical or mental withdrawal symptoms, then they can contact a healthcare provider. These serious symptoms will be recorded in the EDC system.

**Ethics and dissemination**

Ethical approval was obtained from the ethics committee of Sir Run Run Shaw Hospital, an affiliate of Zhejiang University, Medical College (number: 20200129-33). The trial will be performed in accordance with the Declaration of Helsinki. Informed consent will be obtained from all the participants. Each participant will be explained with regards to the purpose, procedures and measurements, potential risks and benefits of the trial before recruitment. All the obtained data will be deidentified and cannot be linked back to the participants. The Unique Identifiers (UIDs) will be automatically deleted from the users’ devices at the end of the trial. The study data collection will start from January 2021. Only the principal investigator and researcher assistants have access to the study data. The primary and secondary analyses will be started immediately after completion of the trial. The results of the trial will be published in the conference proceedings and peer-reviewed journals according to the Consolidated Standards of Reporting Trials statement.
DISCUSSION

This trial will investigate the effectiveness of smartphone apps developed for smoking cessation in China, with the primary outcome of biologically verified continuous smoking abstinence at 6 months, as recommended by the SRNT 26 and the ‘Russell Standard’. 27 To our knowledge, this will be the first RCT to evaluate the efficacy of smoking cessation intervention by using an app based on the cognitive behavioural principles and empirical evidence in Mainland China.

The strength of this study is that it is the first theoretical frameworks guided (CBT), and evidence-based (on the basis of clinical experience, scientific or empirical evidence of smoking cessation) RCT that adheres to the guideline recommendations to evaluate the smoking cessation app in Mainland China. In an era where the prevalence of smartphones and their associated apps have been shown, the trend of exploring and applying these apps in promoting health behaviours such as smoking cessation has become important and inevitable. Smartphone-based smoking cessation apps were easily accessible, highly tailored, intensive but flexible and inexpensive or free of charge. If this app programme is proved to be effective (the funder will further provide financial support to run the app), it opens up a huge potential use in China, with a country of more than 300 million smokers, at the population level, and the impact of public health could be enormous.

The weaknesses of this study include abstinence, which will be mainly measured via a self-report through the app questionnaires, and this is less rigorous than a biochemically verified abstinence. Thus, biological verification for those who have self-reported continuous smoking abstinence at 6 months should be provided and chose this outcome as the primary outcome in this study. In terms of biochemically verified abstinence, participants will receive urine cotinine test by mail (the test results will be confirmed by video call). In order to assess the efficacy of only CBT-based smoking cessation interventions by app, another limitation will be that no first-line medications will be provided to the participants. Furthermore, the participants in this study might be more likely motivated than other smokers, and they might search for other smoking cessation apps or any other available smoking cessation services during the trial. However, the resources that are designed to help people quit smoking and remain smoke-free are extremely limited in China. 28 Finally, participants may be more likely to drop out from the trial, as this smoking cessation app programme will only provide online services. Thus, participants should be encouraged to contact us in case of any question. Also, an ITT approach will be applied to increase the precision and reduce the bias by missing data from loss to follow-up participants.

Text messaging on mobile phones has already been efficacious in helping people to quit smoking in China. 29 Smartphones offer the possibility of supplementing text message-based interventions with computer programmes that can weave together with expert systems, multimedia (eg, music, audio or video), and the internet (eg, WeChat, Weibo or other social networking sites). However, research on content analysis revealed that higher star ratings showed association with better adherence to the 5 As. But most of the currently available smartphone apps for smoking cessation are not particularly ‘smart’, as they have low levels of adherence to the guidelines. 30,31 This new app has been designed and will be revised according to the evidence on smoking cessation principles. This trial assists in building our understanding of how this CBT-based smoking cessation app can be effective in helping people to quit smoking in China.

In summary, this is the first RCT to evaluate the efficacy of CBT-based cigarette smoking cessation smartphone app in China. If this is approved to be effective, then its use will help bridge the huge gap between the unmet needs and the demands for smoking cessation in China at population-based level.

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Competing interests Dr Liao received funding from Johnson & Johnson pharmaceutical company for the study. Dr Tang has no potential conflicts of interest to declare.

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