Abstract

Background: Lifestyle modification (diet, exercise, and behavioral interventions) is the first-line treatment for polycystic ovary syndrome (PCOS). The benefits of face-to-face lifestyle modification intervention in a short time have been demonstrated. However, few studies have investigated the mobile technology effects on lifestyle modification in PCOS. Therefore, we examined the effect of transtheoretical model-based mobile health application intervention program for PCOS.

Methods: A randomised controlled, single-blind trial, was carried out from October 2018 to March 2019, which included 122 participants recruited from gynecology outpatient clinics of affiliated Hospital of Zunyi Medical University in Guizhou. The study participants were randomised into intervention (n = 61) and control groups (n = 61). Participants in the intervention group undertook a TTM-based mobile health application program in addition to routine care, and participants in the control group received only routine care.

Results: Fifty-one participants in the intervention group and 49 in the control group completed the study. Compared to the control group, participants in the intervention group showed statistically significant decrease for BMI (P < 0.05), WC (P < 0.05), SAS (P < 0.05), and SDS (P < 0.05) scores at 6-month and 12-month, respectively. Behavior stage change of exercise and diet among participants with PCOS was significant at 6 months ($\chi^2 = 43.032$, P < 0.05) and 12th months ($\chi^2 = 49.574$, P < 0.05) between the intervention and control groups.

Conclusions: This study showed that the TTM-based mobile health application program can decrease BMI, WC, anxiety, and depression, and improve exercise and diet adherence in patients with PCOS in the long term. The TTM-based mobile health application program can be applied for lifestyle modification in women with PCOS.

Trial registration This study was approved by the ethics committee NO.[2019]1-028 in March 2018 and was registered at the Chinese Clinical Trial Registry (website: www.chictr.org.cn, registry number: ChiCTR2000034572)

Keywords: Polycystic ovary syndrome, Randomised control trial, Mobile health application, Transtheoretical model

Plain Language summary

Poor adherence of life management in patients with PCOS and many then go on to adherence poor health outcomes as a result. There is an urgent need for new approach to enhance to life management compliance for patients with PCOS. TTM is a common behavior change model, which can effectively promote the behavior change of patients, but it has not been used in the life management of PCOS patients. Meanwhile, the traditional life management of PCOS
is mainly face-to-face intervention, few studies have investigated the mobile technology effects on lifestyle modification in PCOS. This study aimed to evaluate the long-term effects of TTM-based mobile technology (for lifestyle modification) on the maintenance of health-related behavior changes among women with PCOS. This study confirmed that the TTM-based mobile health application program can decrease BMI, WC, anxiety, and depression, and improve exercise and diet adherence in patients with PCOS in the long term. Our study conducted TTM-based mobile health application program can provide a reference for other researchers or research organizations.

**Introduction**

Polycystic ovary syndrome (PCOS) is the most common endocrine disorder among women of reproductive age; its prevalence varies from 5 to 20% [1, 2]. Women with PCOS experience a variety of symptoms including polycystic ovaries, hyperandrogenism, and menstrual disturbances; more than 80% of women with PCOS are obese or overweight [3, 4]. Women with PCOS might experience complications associated with their reproductive system in addition to a range of metabolic and psychological consequences (depression, anxiety and lower health-related quality of life) [5], including increased risk of type 2 diabetes (T2D), glucose intolerance, and metabolic syndrome [6]. These physiological symptoms positively associated with anxiety and depression are negatively associated with quality of life [7–9]. Evidence-based guidelines recommend lifestyle modification interventions (including, diet, exercise, and behavioral changes) to be the first-line treatment for PCOS, irrespective of the presenting symptoms [10]. Several studies have demonstrated the short-term effectiveness of face-to-face lifestyle modification interventions [11, 12]. However, poor adherence to lifestyle modification remains a concern, especially in the long term [13, 14].

The transtheoretical model (TTM) (also known as the Stages of Change Theory) was initially developed on the basis of findings about individuals who smoke [15]. This model suggests that behavioral change is a dynamic process, comprising the following five stages [16]: (1) pre-contemplation, which refers to the intention to take action in the following 6 months; (2) contemplation, which refers to the intention to engage in healthy behaviors within the following 6 months; (3) preparation, which refers to the readiness to take action within 1 month; (4) action, which refers to behavior change over the past 6 months; and (5) maintenance; which refers to the sustenance of the behavior change for more than 6 months. Several theories have been used to formulate lifestyle modification interventions, focusing on adherence to behavior change; these theories include the Self-Efficacy Theory, Planned Behavior Theory, and Social Learning Theory. However, intervention studies have mostly used the TTM to improve lifestyle modification adherence owing to its effectiveness in the formulation of intervention strategies corresponding to individual characteristics [17]. Several studies have demonstrated the positive effects of the TTM in the promotion and maintenance of health-related behavior changes among patients with knee osteoarthritis, T2D, obesity, ostomy, etc. [17–19]. However, to the best of our knowledge, few studies have used the TTM to develop and carry out interventions for women with PCOS.

Additionally, it is important to note that smartphone ownership rates in China are considerably high; market research conducted in 2021 estimated that there were 1.07 billion smartphone users in China with the highest proportion of netizens being between the ages of 20 and 39 years [20]. This exponential increase in mobile phone usage in developing countries has been accompanied by rapid developments in mobile health technology. Moreover, studies have suggested that the use of mobile technology for health promotion might be effective in improving long-term health-related outcomes [21, 22]. Mobile technology is a promising tool for intervention pertaining to non-communicable diseases in low-resource settings and developing countries [23–25]. Providentially, mobile transfer of health-related information is an action item in the World Health Organization’s Global Action Plan (2013–2020) for developing countries [26].

As women with PCOS experience symptoms at a young age, there is potential to test the feasibility and effectiveness of mobile health applications for improving health-related behavior among these women. However, limited research has been conducted on the same; to the best of our knowledge, only two studies so far have focused on the design of a personalized mobile tool called AskPCOS. Research indicates that currently available applications are unlikely to meet their information needs. Additionally, no study has investigated the effectiveness of mobile technology in the interventions for PCOS [27, 28].

In light of the aforementioned arguments, it is necessary to investigate the effectiveness of TTM-based mobile technology for lifestyle modification among women with PCOS. Thus, this study aimed to evaluate
the long-term effects of TTM-based mobile technology (for lifestyle modification) on the maintenance of health-related behavior changes among women with PCOS.

Materials and methods
Design, setting, and participants
This study was performed using a single-blind randomized controlled trial. Participants were recruited from outpatient gynecology clinics—of the hospital affiliated with Zunyi Medical University in Zunyi—from October 2018 to March 2019. Zunyi is the second largest city in the Guizhou province of China with a population of 6.3 million people, covering an area of 30,762 km². It is located in the neighborhood of Luzhou (in the Sichuan Province) and bordered by Chongqing in the north.

Participants included in the study (1) fulfilled at least two of the three Rotterdam Criteria for the diagnosis of PCOS (including oligo-ovulation or anovulation, hyperandrogenism, and polycystic ovaries that were confirmed via ultrasound) [29]; (2) were aged 18 years or above and had a BMI that was greater than or equal to 25 kg/m²; (3) had access to a smartphone; and (4) had a sedentary lifestyle. Participants who were pregnant, had severe disabilities which prevented them from taking care of themselves, or served as participants in other intervention programs for PCOS were excluded from the study.

Sample size and randomization
A total of 122 participants (61 participants per group) were recruited before the intervention according to inclusion and exclusion criteria. The sample size adequately met the parametric test assumptions, yielding an impact size of 0.30, a power of 90% with 5% margin of error, and allowing an attrition rate of 10%.

An independent researcher prepared the randomization sequence on the basis of a computer-generated block randomization list for 1:1 allocation of participants into two groups (intervention and control groups) with a block size of four. Following the baseline data collection process, participants were randomly assigned to the groups by a researcher who was blinded to the study design.

Ethical considerations
This study was approved by the institutional review board (IRB) of the hospital affiliated with Zunyi Medical University (Approval no.: 2019/1-028) prior to the commencement of data collection. All participants provided written informed consent for their participation in the study. Furthermore, no adverse events occurred.

Procedures
Gynecologists identified potentially eligible participants who visited the outpatient gynecology clinics. If the participants fulfilled the inclusion criteria, they were acquainted with the purpose, procedures, and benefits of the study. Data were then collected from participants who provided written informed consent for their participation in the study. Following the baseline data collection process, participants in the intervention group were trained by the researchers to use the mobile health application. Participants attended the outpatient clinic for standardized screening, which was conducted by two researchers who were blinded to group allocation; the measurement time points were as follows: baseline, 6 months, and 12 months. The differences in outcome measures between the intervention and control groups were evaluated across the measurement time points to check for the long-term effectiveness of the TTM-based mobile health application.

Intervention group
The participants in the intervention group were required to use the TTM-based mobile health application called Home of PCOS, comprising two modules. A multi-disciplinary team—including an obstetrics and gynecology professor, a nursing science professor, a nutritionist, a psychologist, and a software engineer—designed the application. The application was published in collaboration with a software company.

The first module—titled ‘Assessment of my current behavioral stage’—of the application required participants to answer a question pertaining to their dietary behaviors and exercise by selecting one of the three alternatives. Based on their responses, participants were divided into the pre-action, action, and maintenance subgroups. Participants were re-assessed for their stage of behavior change every month; accordingly, they were assigned to the relevant subgroup. The second module—titled ‘What I need to do And how?’—comprised three separate links for the pre-action, action, and maintenance subgroups. Participants were assigned a link based on the assessment of their current behavioral stage; the link comprised specific interventions for the participants.

Pre-action subgroup: This intervention subgroup included three steps (or pages). The first page—titled Knowledge and information about PCOS—contained recorded lectures and resource pages. The second page—titled ‘How to prepare for lifestyle change?’—was the provision of knowledge and information to the participants who were preparing for lifestyle changes. The third page—titled ‘You need to change your lifestyle’—was an online chat room.
Action subgroup: This intervention subgroup also included three steps (or pages). The first page—titled ‘Self-monitoring’—required participants to record their exercise and diet. The second page—titled ‘Peer support’—was an online chat room. The third page—titled ‘Expert consultation’—was an online chat room.

Maintenance subgroup: This intervention subgroup included two steps (or pages). The first page—titled ‘Keep up your adherence to lifestyle modification’—Provided knowledge and information. The second page—titled ‘Expert consultation’—was an online chat room.

Table 1 shows a detail description of all intervention components.

Control group
Participants in the control group received routine care; that is, during their visits to the outpatient gynecology clinics, gynecologists gave them advice regarding the maintenance of physical activity and well-balanced diet. The gynecologist provided the same advice to participants in the intervention group.

Outcome measures
Sociodemographic characteristics
Demographic characteristics were measured using the personal information form, which was developed by the researcher and comprised seven questions pertaining to gender, ethnicity, education level, living residence, marital status, occupation, and years of PCOS.

Body mass index
The researchers used an electronic scale to measure the participants’ weight. Participants’ height was measured as the length from the top of their head to their sole while they stood in the Frankfort plane (that is, the back of the skull, shoulders, pelvis, and heels touched the same horizontal plane when they stood at attention). It was ensured that the participants were assessed barefoot and in light clothes. The body mass index (BMI) was calculated as weight/height² (kg/m²) [30].

Waist circumference
Waist circumference (WC) of the individuals was measured over their underwear—after a mild expiration while standing—by placing the measuring tape between the edge of the lower costal margin and iliac crest [30]. This measurement was conducted twice per participant and the mean of the two values was used for data analysis.

Anxiety
Anxiety was measured using Zung’s Self-Rating Anxiety Scale (SAS), which comprises 20 items that are rated on a 4-point Likert scale, ranging from 1 (“none or a little of the time”) to 4 (“most or all of the time”); total scale scores range from 20 to 80. Cronbach’s alpha for the SAS

| Stage of change | Intervention module | Detail strategy |
|-----------------|---------------------|----------------|
| Pre-action (three steps) | ① Knowledge and information about PCOS | Contained recorded lectures and resource pages (such as ‘Basic knowledge about PCOS’ and ‘The benefits of a healthy lifestyle for patients with PCOS’) that provided knowledge and information about PCOS |
| | ② How to prepare for lifestyle change | Provided of knowledge and information to the participants who were preparing for lifestyle changes. It contained eight recorded lectures on topics such as home-based exercise programs (including exercise types, duration, and timings), diet and nutrition in PCOS, eating patterns (focusing on the importance of frequency and regularity), and the development of suitable exercise and diet plans |
| | ③ You need to change your lifestyle | In an online chat room: enable one-on-one communication between each participant and the professionally trained researcher, a couple of times; the researcher applied the motivation interview technology to trigger participants’ motivation for modifying their lifestyle |
| Action (three steps) | ① Self-monitoring | Required participants to record their exercise routine (including exercise type, duration, time, and intensity) per week and diet (including daily meals and daily energy intake) per day |
| | ② Peer support | Included a chat room where participants could share their exercise and diet-control experiences with each other |
| | ③ Expert consultation | Included a chat room where participants could consult experts and seek strategies to overcome barriers to lifestyle modification |
| Maintain (two steps) | ① Keep up your adherence to lifestyle modification | Provided knowledge and information to help participants in sustaining their adherence to lifestyle modification |
| | ② Expert consultation | Included a chat room where participants could consult experts and seek strategies to overcome barriers to lifestyle modification |
was 0.82 [31]. Zung recommended the conversion of raw scores to index scores (ranging from 25 to 100) by multiplying the raw score with 1.25. The SAS scores were correlated with the Taylor Manifest Anxiety Scale scores \((r = 0.30)\) [32]. Thus, the SAS is a psychometrically sound measure of anxiety with good sensitivity, specificity, and the capacity to discriminate between anxiety and other psychiatric disorders [2].

**Depression**
Depression status was assessed using Zung’s Self-Rating Depression Scale (SDS), which comprises 20 items that are rated on a 4-point Likert scale, ranging from 1 (“none or a little of the time”) to 4 (“most or all of the time”); total scale scores range from 20 to 80. However, Zung [33] recommended the conversion of raw scores to index scores (ranging from 25 to 100) by multiplying the raw score with 1.25.

**Behavior change stage**
The behavior change stage for exercise and diet was assessed using the following question: “Are you exercising at least thrice a week (for 150 min per week) at a moderate level and maintaining dietary restraint everyday (for example, consuming a low-fat diet)?” Participants responded by choosing one of the following alternatives, which determined the behavior change stage they were in: (a) No, I have never done this (pre-action stage); (b) Yes, I am; however, I have been doing this for less than 6 months (action stage); and (c) Yes, I have been doing this for more than 6 months (maintenance stage).

**Statistical analysis**
The SPSS Software (version 18.0) was used for data analysis. Normally distributed data for continuous variables were analyzed using means and standard deviations; whereas, non-normally distributed data were expressed as percentages. Independent sample t-tests were conducted to analyze the statistical significance of the differences in the means of continuous variables between the intervention and control groups. A repeated measures ANOVA was conducted to determine the statistical significance of the effects of the intervention within and between the groups. The Mann–Whitney U and Chi-square tests were conducted to evaluate the statistical significance of differences in the general data (ethnicity, education level, living residence, marital status, occupation, and years of PCOS) between the two groups.

**Results**
Of the 122 participants enrolled in the trial between February 2020 and November 2021, 100 participants completed the study with 51 participants in the intervention group and 49 participants in the control group. Figure 1 shows the participant flow throughout the study. There were no statistically significant differences in the
sociodemographic characteristics between participants in the intervention and the control groups (P > 0.05; Table 2).

Furthermore, there were no statistically significant differences in the BMI (P = 0.38), WC (P = 0.43), SAS scores (P = 0.223), and SDS scores (P = 0.621) between the intervention and control groups at the baseline (Table 3). However, the group-by-time interaction effects on the BMI, WC, SAS scores, and SDS scores were statistically significant, indicating significant group differences on different occasions (P < 0.05). The BMI of participants in the intervention group significantly decreased from the baseline (M = 25.99, SD = 3.87) to the 12th month (M = 22.63, SD = 1.97). Similarly, the WC of participants in the intervention group significantly decreased from the baseline (M = 86.22, SD = 9.91) to the 12th month (M = 80.50, SD = 6.94). However, there were no statistically significant differences in the BMI and WC at different measurement time points among participants in the control group.

In addition, the SAS scores of participants in the intervention group significantly decreased from 46.97 (SD = 6.53) at the baseline to 42.12 (SD = 9.69) at the 12th month (P = 0.035). The SDS scores of participants in the intervention group significantly decreased from the baseline (M = 51.55, SD = 6.78) to the 12th month (M = 40.57, SD = 5.41; P < 0.001). In contrast, there were no statistically significant differences in the SAS and SDS scores at different measurement time points among participants in the control group.

There was a significant difference in the behavior change stage for exercise and diet between the intervention and control groups at the 6th month (c^2 = 43.032, P < 0.05); that is, 82.4% of the participants in the intervention group were in the action stage of behavior change, whereas, there was no significant change in behavior stage from the baseline to the 6th month (P > 0.05) among participants in the control group (Fig. 2).

Furthermore, there was a significant difference in the behavior change stage for exercise and diet between the intervention and control groups at the 12th month (c^2 = 49.574, P < 0.05). In contrast, there was no significant change in the behavior stage among participants in the control group at the 12th month (P > 0.05). Interestingly, 33.3% of the participants in the intervention group had transitioned to the action stage, and 66.7% of the participants had transitioned to the maintenance stage; whereas only 10.2% of participants in the control group had completed the pre-action and action stages, and entered the maintenance stage.

### Table 2: Comparison of sociodemographic characteristics of the recruited participants at baseline

| Variable | Categories | Intervention group (N = 51) | Control group (N = 49) | χ²/t | P value |
|----------|------------|-----------------------------|------------------------|------|--------|
| Age (x ± s) | 24.72 ± 4.20 | 24.94 ± 4.31 | 0.37* | 0.71 |
| Ethnic group [N(%)] | Han-nationality | 31 | 27 | 0.331 ▲ | 0.565 |
| Ethnic minority | 20 | 22 |
| Living residence [N(%)] | City | 22 | 21 | 0.001 ▲ | 0.997 |
| Countryside | 29 | 28 |
| Marital status [N(%)] | Single | 29 | 27 | -0.205 ▼ | 0.838 |
| Married | 19 | 17 |
| Widowed/divorced | 3 | 5 |
| Education [N(%)] | Middle school | 10 | 9 | 0.366 ▲ | 0.947 |
| High school | 12 | 14 |
| junior college | 5 | 4 |
| College | 24 | 22 |
| Occupation [N(%)] | Employed | 13 | 14 | 0.856 ▲ | 0.836 |
| Unemployed | 12 | 8 |
| Student | 18 | 18 |
| Other | 8 | 9 |
| Years of PCOS [N(%)] | < 1 years | 28 | 26 | -0.38 ▼ | 0.62 |
| 1–3 years | 14 | 12 |
| 4–6 years | 4 | 4 |
| > 7 years | 5 | 7 |

* t; ▲ χ²; ▼ Mean-Whitney U
Discussion

A review of the extant literature suggests that the findings pertaining to the feasibility and effectiveness of mobile health interventions are inconsistent. A previous study comparing the benefits of mobile health applications with routine care among people with chronic obstructive pulmonary disease found no statistically significant differences in improving the health-related quality of life; whereas few studies have highlighted the effectiveness of mobile-application based health interventions.

The current study was one of the few studies that examined the effectiveness of a TTM-based mobile health intervention program (using the mobile application called Home of PCOS) on the maintenance of self-managed health-related behavior change among women with PCOS. BMI and WC are important clinical indices that are associated with PCOS symptoms. A recent systematic review reported that dietary modifications alone were associated with a significant decrease in the BMI [34]. Another systematic review showed a statistically significant decrease in the BMI and WC as a consequence of short-duration (less than 12 weeks) and supervised aerobic exercise-based interventions with or without diet combined [35]. On the contrary, the current study found that the maintenance of self-managed health-related behavior change—including exercise and diet—led to a significant decrease in the BMI and WC over time (at both 6 and 12 months), highlighting the efficacy of the combination of routine care with the use of Home of PCOS (intervention group) as opposed to only receiving routine care (control group). These findings could be attributed to the TTM-based mechanisms.

### Table 3 BMI, WC, SAS and SDS of change at baseline, 6-month and 12-month post-intervention among the PCOS patients

| Measured parameter | Baseline | 6-month | 12-month | F(time*group) | P value |
|--------------------|----------|---------|----------|---------------|---------|
| BMI                |          |         |          |               |         |
| Intervention group | 25.99±3.87| 23.40±1.69| 22.63±1.97| 12.8         | 0.00**  |
| Control group      | 25.25±3.95| 25.36±3.06| 24.86±2.65|               |         |
| t-value            | 0.89     | −3.7    | 6.55     |               |         |
| P-value            | 0.38     | 0.00**  | 0.00**   |               |         |
| WC                 |          |         |          |               |         |
| Intervention group | 86.22±9.91| 81.54±8.54| 80.50±6.94| 2.44         | 0.04*   |
| Control group      | 87.64±6.59| 86.75±5.52| 85.44±7.95|               |         |
| t-value            | −0.79    | −3.44   | −3.14    |               |         |
| P-value            | 0.43     | 0.00**  | 0.00**   |               |         |
| SAS                |          |         |          |               |         |
| Intervention group | 46.97±6.53| 42.91±6.65| 42.12±9.69| 0.91         | 0.016*  |
| Control group      | 49.02±7.05| 47.83±6.27| 46.46±8.91|               |         |
| t-value            | −1.23    | −3.49   | −2.14    |               |         |
| P-value            | 0.223    | 0.001** | 0.035*   |               |         |
| SDS                |          |         |          |               |         |
| Intervention group | 51.55±6.78| 46.83±5.36| 40.57±5.41| 8.41         | 0.000***|
| Control group      | 52.29±6.86| 50.57±5.42| 49.24±7.14|               |         |
| t-value            | 0.5      | −2.72   | −6.24    |               |         |
| P-value            | 0.621    | 0.008** | 0.000*** |               |         |

F(time*group): The changing trend of measurement indicators in intervention group and control group at different intervention time points.

BMI body mass index, WC Waist circumference, SAS Self Rating Anxiety Scale, SDS Self Rating Depression Scale.

*P < 0.05; **P < 0.01; ***P < 0.001

### Fig. 2 Behavior stage change of exercise and diet among patients with PCOS at the 6th month
underlying the mobile health application in the current study that facilitated the initiation and management of behavior change among participants in the intervention group. During the action stage, participants used strategies such as self-monitoring and peer support—through the mobile application—which helped them to maintain health-related behaviors (Fig. 3).

Women with PCOS experience higher levels of anxiety and depressive symptoms as compared to women without PCOS [36, 37], prompting international guidelines to recommend the management of anxiety and depressive symptoms among women with PCOS [38]. Some studies have suggested that lifestyle management could result in significantly lower levels of depression [39, 40]. However, a 20-week long diet and exercise intervention study found that low level of depressive symptoms among participants in the intervention group could only be sustained during the first 10 weeks [41]. Furthermore, two diet-induced weight loss studies also found short-term improvement in psychological outcomes; however, this improvement could not be sustained in the long-term [42, 43]. Nevertheless, the current study found significant improvements in anxiety and depressive symptoms among participants in the intervention group at 6 months; the improvements were sustained at 12 months. The aforementioned findings can be attributed to the following factors. First, significant weight loss among women with PCOS has been found to have positive effects on psychological outcomes [44]. Second, lack of information for the management of PCOS might lead to anxiety and depressive symptoms among women with PCOS [45]. The present study facilitated the provision of PCOS-related information to participants in the intervention group via the mobile health application. This might have had positive implications on the participants’ psychological health.

Maintaining health-related behavior changes in the long term is challenging for women with PCOS. Many traditional face-to-face interventions have reported poor adherence to lifestyle modification. A systematic review reported [6] that only 50% of PCOS patients persisted in lifestyle modification during lifestyle intervention in the short term (less than 12 months). Another study involving a 12-month lifestyle intervention for women with PCOS reported 54% adherence to lifestyle modification [46]. To the best of our knowledge, this is the first study to have investigated the long-term effectiveness of a TTM-based mobile health application on health-related behavior changes (including exercise and diet) among patients with PCOS. The positive effects of the TTM-based mobile health application on the stages of behavior change could be attributed to several factors. First, the transition to the action stage primarily depends on their knowledge, skills, and motivation. The TTM-based mobile health application provided knowledge, and strategies for lifestyle modification, in addition to motivating the participants to adhere to their lifestyle changes. Second, most people who take action to change their behaviors are not successful in maintaining it in the long term [47]. In the current study, participants were assessed for their behavior change stage for diet and exercise every month; accordingly, the mobile health application provided corresponding intervention strategies. The intervention strategies for improving health-related behavior were flexible; this could have prevented the regression to the previous behavior change stages. Third, mental health outcomes such as depression and anxiety negatively influence trial adherence [48]. In this study, we found significant improvements in anxiety and depressive symptoms at 6 months among participants in the intervention group; these improvements were sustained at 12 months. This might have positively influenced the adherence to healthy behaviors.

It is important to acknowledge the limitations of this study. First, participants were recruited from one general hospital; this might have limited the generalizability of the findings. Second, the study assessed self-reported adherence to the health-related behavior changes; these changes were not monitored by the researchers. Furthermore, recall bias could have reduced the accuracy of the findings. Third, the time span for follow-up regarding health-related behavior change was short (12 months). Future studies with longer duration of intervention, and carry out more TTM-based mobile health applications multicenter RCT research are required to confirm the validity of our findings.
Conclusion

This study showed that the TTM-based mobile health application—Home of PCOS—had significant long-term effectiveness in reducing BMI, WC, anxiety, and depressive symptoms and improving adherence to health-related behavior changes among women with PCOS. The application can be used in addition to the provision of routine care to facilitate adherence to lifestyle modification among women with PCOS.

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Author contributions

LHW drafted the work and revised it critically for important intellectual content. YL and HT participated in the work’s acquisition, analysis, or interpretation of data. SH carried out the conception, design of the work and the final approval of the version to be published and agreed to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. All authors read and approved the final manuscript.

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Availability of data and materials

The dataset used during the current study is available from the corresponding author on reasonable request.

Declarations

Ethics approval and consent to participate

The research protocol was approved by the Institutional Ethical Review Board at the Affiliated Hospital of Zunyi Medical University. Written informed consent was obtained from all participants.

Consent for publication

All participants signed informed consent forms.

Competing interests

The authors declare that they have no competing interests.

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