Classroom-based cognitive behaviour therapy (FRIENDS): a cluster randomised controlled trial to Prevent Anxiety in Children through Education in Schools (PACES)

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Classroom-based cognitive behaviour therapy (FRIENDS): a cluster randomised controlled trial to Prevent Anxiety in Children through Education in Schools (PACES)

Paul Stallard, Elena Skryabina, Gordon Taylor, Rhiannon Phillips, Harry Daniels, Rob Anderson, Neil Simpson

Summary

Background Anxiety in children is common, impairs everyday functioning, and increases the risk of severe mental health disorders in adulthood. We investigated the effect of a classroom-based cognitive behaviour therapy prevention programme (FRIENDS) on anxiety symptoms in children.

Methods Preventing Anxiety in Children through Education in Schools (PACES) is a three-group parallel cluster randomised controlled trial. Interventions were given between September, 2011, and July, 2012, with schools as the unit of allocation and individual participants as the unit of analysis. We enrolled state-funded junior schools in southwest England. We sent information to all eligible schools (state-funded junior schools in southwest England) inviting them to enrol in the study. School year groups were assigned by computer-generated randomisation (1:1:1) to receive either school-led FRIENDS (led by teacher or school staff member), health-led FRIENDS (led by two trained health facilitators), or usual school provision. Children were not masked to treatment allocation. The allocated programme was given to all students (aged 9–10 years) in the school year (ie, universal delivery) as part of the school curriculum as nine, 60 min weekly sessions. Outcomes were collected by self-completed questionnaire administered by researchers masked to allocation. Primary outcome was symptoms of anxiety and low mood at 12 months assessed by the Revised Child Anxiety and Depression Scale (RCADS 30). Analyses were intention to treat and accounted for the clustered nature of the design. The study is registered, number ISRCTN23563048.

Findings 45 schools were enrolled: 14 (n=497 children) were randomly assigned to school-led FRIENDS, 14 (n=509) to health-led FRIENDS, and 12 (n=442) to usual school provision. 1257 (92%) children completed 12 month assessments (449 in health-led FRIENDS, 436 in school-led FRIENDS, and 372 in usual school provision). We recorded a difference at 12 months in adjusted mean child-reported RCADS scores for health-led versus school-led FRIENDS (19·49 [SD 14·81] vs 22·86 [15·24]; adjusted difference −3·91, 95% CI −6·48 to −1·35; p=0·0004) and health-led FRIENDS versus usual school provision (19·49 [14·81] vs 22·48 [15·74]; −2·66, −5·22 to −0·09; p=0·043). We noted no differences in parent or teacher ratings. Training teachers to deliver mental health programmes was not as effective as delivery by health professionals.

Interpretation Universally delivered anxiety prevention programmes can be effective when used in schools. However, programme effectiveness varies depending on who delivers them.

Funding National Institute for Health Research Public Health Research Programme.

Introduction Anxiety disorders affect 10% of children by the age of 16 years. They significantly impair everyday functioning, often persist into adulthood, and increase the risk of other psychiatric disorders in adolescence and young adulthood. The associated health-related burden, and economic and societal costs are large, and the need to improve the mental health of children is being increasingly recognised as a global priority.

Effective psychological interventions, especially cognitive behaviour therapy (CBT), are available for children with anxiety disorders. However, comparatively few children with anxiety disorders are identified and referred for treatment. The poor reach and availability of traditional treatment services has led to interest in more proactive preventive approaches with schools offering a convenient and natural location to deliver such programmes. Findings of systematic reviews show that universal and targeted prevention programmes are often based on cognitive behaviour therapy. Data from reviews suggest that cognitive behaviour therapy prevention programmes can be effective, although research methods are poor, adequately powered implementation trials are scarce, results are inconsistent, and effect sizes vary greatly. Most studies report overall changes in symptoms, and the preventive benefits for less symptomatic participants have seldom been reported.

Prevention programmes can be universally provided to all of an identified population, or targeted towards those at risk of developing a disorder or showing early signs of a disorder, or a combination of both approaches. Universal programmes have good reach, avoid the need for screening, are less stigmatising, and offer the potential to enhance mental health and reduce present symptoms.
Targeted programmes focus scarce resources on individuals with greatest needs, and usually achieve larger treatment effects.\(^1\)\(^2\)\(^3\)

The effect of the intervention leader (health vs school professional) has important implications for the method of delivery and sustainability of an intervention but has been directly investigated in only one study.\(^1\)\(^9\) Barrett and Turner noted that a universal anxiety prevention programme (FRIENDS; panel 1) was equally effective in the reduction of symptoms of anxiety in children aged 10–12 years when given by a psychologist or teacher. However, systematic reviews have reached different conclusions about who is most effective at delivering these programmes.\(^1\)\(^5\)\(^6\)

Before anxiety prevention programmes can be endorsed and widely provided, independent implementation trials are needed to measure effectiveness and cost-effectiveness when provided under real-life conditions and to establish the effect of the intervention leader on outcome.

We undertook a pragmatic assessment of the effectiveness of a classroom-based anxiety prevention programme (FRIENDS\(^2\)) universally delivered by health and school professionals to school years 4 and 5 (children aged 9–10 years) in UK junior schools.

**Methods**

**Study design and participants**

We did this three-group parallel cluster randomised controlled trial between September, 2011, and July, 2012, with school as the unit of allocation and individual participants as the unit of analysis.\(^2\)\(^0\) A project information sheet and trial enrolment form was sent to all primary schools in Bath and northeast Somerset, Swindon Borough, and Wiltshire within a 50 mile radius of the University of Bath, UK (n=268). Eligible schools were state-funded junior schools in three Local Education Authorities in southwest England. Such junior schools are mainstream government-funded schools that are attended by 94·5% of children aged 5–10 years in the UK. All children aged 9–10 years (years 4 and 5) in participating schools were eligible, unless they were not attending school (eg, because of long-term sickness or excluded from school) or did not participate in Personal Social and Health Education (PSHE) lessons for religious or other reasons. The allocated intervention was given to all participants in the school year (ie, universal delivery) as part of the PSHE curriculum. The trial protocol is online.

Participation required written consent from the school head teacher, parents’ not opting their child out of the study, and signed assent from the child. The study was approved by the University of Bath, Department for Health Research Ethics Committee.

**Randomisation and masking**

Once all schools had been enrolled, we randomly assigned year groups (1:1:1) to school-led FRIENDS, health-led FRIENDS, or usual school provision. Randomisation was undertaken at school and not the class level to avoid possible contamination within schools. Trial groups were balanced with respect to key characteristics by calculating an imbalance statistic for a large random sample of possible allocation sequences.\(^2\)\(^3\) Children were not masked to treatment allocation. Outcomes were collected by self-completed questionnaire administered by researchers masked to allocation. Group allocation was kept in a separate password-protected database. Researchers who analysed data were also masked to allocation—trial groups were numerically coded and data analysis undertaken masked to which code related to each trial group. The variables used for balancing were school size, number of students and classes, number of mixed classes, level of educational attainment, and preferred timetabling. A statistician with no other involvement in

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**Panel 1: Acronym for the FRIENDS process**

- F: Feelings
- R: Remember to relax
- I: I can do it. I can try my best
- E: Explore solutions and coping step plans
- N: Now reward yourself. You’ve done your best
- S: Smile. Stay calm for life

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**For more on the trial protocol see** [http://www.trialsjournal.com/content/13/1/227/abstract](http://www.trialsjournal.com/content/13/1/227/abstract)

**Figure: Trial profile**
the study randomly selected one sequence from a subset with the most desirable balance properties.

Procedures
Interventions were delivered in the academic year September, 2011, to July, 2012. The anxiety prevention programme we assessed (FRIENDS) is a manualised cognitive behaviour therapy intervention that has been identified as effective. The programme was developed to be delivered in schools and provided to whole classes of children. FRIENDS is based on the principles of CBT and develops skills to counter the cognitive, emotional, and behavioural aspects of anxiety. Children develop emotional awareness and regulation skills, to identify and replace cognitions that increase anxiety with more balanced and functional ways of thinking and to develop problem-solving skills to confront and cope with situations and events that provoke anxiety. The intervention trialled in this study consisted of nine, 60 min weekly sessions delivered to whole classes of children. Children had their own workbook consisting of nine, 60 min weekly sessions delivered to provoke anxiety. The intervention trialled in this study randomly selected one sequence from a subset with the most desirable balance properties.

The intervention trialled in this study consisted of nine, 60 min weekly sessions delivered to whole classes of children. Children had their own workbook and group leaders had a detailed session plan that specified key learning points, objectives, and core activities for each session. The feasibility and viability of delivering FRIENDS in UK schools has previously been established.

In the health-led FRIENDS programme, each session was led by two trained health facilitators working alongside the class teacher. All facilitators had at least an undergraduate university degree in a relevant discipline (eg, social science or education), appropriate professional backgrounds (eg, psychology, nursing, or education), or experience of working with children or young people. Initial 2 day training and supervision every 2 weeks were provided by accredited FRIENDS trainers. Supervision was in a group format and consisted of review of session plans, the underlying cognitive model, class and behaviour management skills, and any interpersonal difficulties or communication problems with the class teacher.

In the school-led FRIENDS programme, sessions were led by a teacher or member of the school staff (eg, teaching assistant) who were trained in delivery of the programme and were supported by two facilitators. School staff attended the same 2 day initial training and were offered ongoing supervision. We assessed treatment fidelity by randomly assessing audiotape recordings of 10% of FRIENDS sessions.

In the usual school provision programme, children participated in the usual PSHE sessions provided by the school. All schools were following a UK National Curriculum programme designed to develop self-awareness, management of feelings, motivation, empathy, and social skills. The sessions were planned and provided solely by the teacher and did not include any external input from the research team.

Outcomes
Child outcomes were collected during class time with self-completed questionnaires administered by researchers at baseline, 6 months, and 12 months. As described in the trial protocol, the primary outcome was symptoms of anxiety and low mood 12 months after baseline as established by the Revised Child Anxiety and Depression Scale (RCADS 30). Secondary outcomes assessed worry (with the Penn State Worry Questionnaire for Children), self-worth and acceptance (with the Rosenberg Self-Esteem Scale), extent of bullying (with the Olweus Bully/Victim Questionnaire), and life satisfaction (with a subjective wellbeing assessment).

<table>
<thead>
<tr>
<th>Health-led FRIENDS (N=489)</th>
<th>School-led FRIENDS (N=472)</th>
<th>Usual PSHE (N=401)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of schools</td>
<td>14</td>
<td>14</td>
</tr>
<tr>
<td>Number of schools with 2+ classes</td>
<td>6</td>
<td>7</td>
</tr>
<tr>
<td>Class size</td>
<td>19·56 (6·56)</td>
<td>18·15 (7·68)</td>
</tr>
<tr>
<td>Missing baseline assessment</td>
<td>3 (0·6%)</td>
<td>10 (2·1%)</td>
</tr>
<tr>
<td>Sex</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Boys</td>
<td>255 (52·1%)</td>
<td>237 (50·2%)</td>
</tr>
<tr>
<td>Girls</td>
<td>234 (47·9%)</td>
<td>235 (49·8%)</td>
</tr>
<tr>
<td>Ethnic origin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>British white</td>
<td>455 (94·2%)</td>
<td>439 (95·2%)</td>
</tr>
<tr>
<td>Non-white</td>
<td>28 (5·8%)</td>
<td>22 (4·8%)</td>
</tr>
<tr>
<td>Living situation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mother and father</td>
<td>347 (71·4%)</td>
<td>315 (68·2%)</td>
</tr>
<tr>
<td>Parent and partner</td>
<td>43 (8·8%)</td>
<td>55 (11·9%)</td>
</tr>
<tr>
<td>Single parent</td>
<td>67 (13·8%)</td>
<td>68 (14·8%)</td>
</tr>
<tr>
<td>Other</td>
<td>29 (6·0%)</td>
<td>24 (5·2%)</td>
</tr>
<tr>
<td>Number of siblings</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>49 (10·1%)</td>
<td>30 (6·5%)</td>
</tr>
<tr>
<td>1</td>
<td>221 (45·5%)</td>
<td>214 (46·5%)</td>
</tr>
<tr>
<td>2</td>
<td>129 (26·5%)</td>
<td>134 (29·1%)</td>
</tr>
<tr>
<td>3 or more</td>
<td>87 (17·9%)</td>
<td>82 (17·8%)</td>
</tr>
<tr>
<td>Family affluence</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low (0–2)</td>
<td>6 (1·5)</td>
<td>11 (2·4)</td>
</tr>
<tr>
<td>Medium (3–5)</td>
<td>142 (29·4)</td>
<td>139 (30·1)</td>
</tr>
<tr>
<td>High (6–8)</td>
<td>331 (69·1)</td>
<td>311 (67·5)</td>
</tr>
<tr>
<td>Child total RCADS</td>
<td>26·24 (15·56)</td>
<td>24·91 (14·32)</td>
</tr>
<tr>
<td>Penn Worry Scale</td>
<td>10·63 (8·14)</td>
<td>10·99 (8·24)</td>
</tr>
<tr>
<td>Self-esteem</td>
<td>18·94 (5·34)</td>
<td>19·43 (5·39)</td>
</tr>
<tr>
<td>Total life satisfaction</td>
<td>14·21 (6·77)</td>
<td>13·32 (5·71)</td>
</tr>
<tr>
<td>Been bullied</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bullied at school</td>
<td>142 (29·3%)</td>
<td>124 (26·8%)</td>
</tr>
<tr>
<td>Not bullied or only once</td>
<td>343 (70·7%)</td>
<td>338 (73·2%)</td>
</tr>
<tr>
<td>Parent-completed assessments</td>
<td>217 (44·4%)</td>
<td>201 (42·6%)</td>
</tr>
<tr>
<td>Total RCADS</td>
<td>12·55 (8·83)</td>
<td>10·99 (8·60)</td>
</tr>
<tr>
<td>Total SDQ</td>
<td>9·09 (6·32)</td>
<td>8·31 (6·28)</td>
</tr>
<tr>
<td>Teacher-reported assessment</td>
<td>487 (99·6%)</td>
<td>466 (98·7%)</td>
</tr>
<tr>
<td>Teacher SDQ impact</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Difficulty</td>
<td>119 (24·4)</td>
<td>125 (26·8)</td>
</tr>
</tbody>
</table>

Data are mean (SD) or number (%) unless otherwise indicated. Family affluence assessed by the Family Affluence Scale. PSHE—personal social and health education. RCADS—Revised Child Anxiety and Depression Scale. SDQ—Strength and Difficulties Questionnaire.

Table 1: Baseline characteristics
Parents completed a behavioural screening questionnaire (Strength and Difficulties Questionnaire [SDQ]) and the parent version of the Revised Child Anxiety and Depression Scale (RCADS-30-P) at baseline, 6 months, and 12 months.

Class teachers completed the impact rating of the Strengths and Difficulties Questionnaire (SDQ) for all children in their class at all three assessment points to assess the presence of an emotional or behavioural problem, chronicity, distress, social impairment, and burden.

### Statistical analysis

We powered the study to detect a difference of 3-6 points in mean RCADS total scores between FRIENDS and usual PSHE. On the basis of an SD of 12 points and an intracluster correlation coefficient of 0-02, 28 pupils per class, 90% consent and 80% retention, effect sizes between 0.28 and 0.30, SDs are detectable with 80% power and 5% two-sided α with 1134-1360 assenting pupils.

We used descriptive statistics to assess balance between the trial groups at baseline. The primary outcome was assessed by intention to treat without imputation. To take...
appropriate account of the hierarchical nature of the data, we used multivariable mixed effects models to compare mean RCADS at 12 months for health-led FRIENDS with school-led FRIENDS and usual school provision, with adjustment for baseline RCADS, sex, and school effects. We repeated these analyses for secondary outcomes. For RCADS we undertook a further planned analysis. We used repeated-measures mixed-effects analysis of variance models to investigate convergence and divergence between trial groups over time. We did preplanned subgroup analyses with interaction terms in the regression models between the randomised group and the baseline variable (low anxiety RCADS 0–48, high anxiety ≥49).

We did sensitivity analyses to assess the potential effect of missing data. Completion rates for all groups at 12 months were high (91·8–92·7%), although non-completers tended to be more symptomatic on our primary outcome measure (RCADS) at baseline (data not shown). With multiple imputation methods, we created 20 datasets and showed that imputation for missing data made no material difference to the overall results. Therefore, the data we present are based on recorded data only.

The study is registered, number ISRCTN23563048.

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

Results
Between September, 2011, and July, 2012, 45 schools were enrolled. 41 consented to participate and were randomly assigned: 14 (n=497 children) to school-led FRIENDS, 14 (n=509) to health-led FRIENDS, and 12 (n=442) to usual school provision. One school from usual school provision withdrew before baseline assessments were undertaken (figure). The remaining 40 schools were representative of the UK in terms of academic attainment according to Department of Education performance tables (ie, the percentage of children achieving key stage 2 level 4 in maths and English; data not shown). However, more children had special educational needs (23·2% vs 17·1%), pupil absence rates were lower (4·4% vs 5·1%), and eligibility for free school meals was lower (12·4% vs 18·2%) in the cohort than the national average.

Of the 1448 eligible participants, 1362 (94%) consented to participate in the study, of whom 1339 (98%) completed baseline assessments. The proportion of boys in the usual school provision group (42%) was lower than in each of the other two trial groups, but otherwise the groups were well balanced (table 1).

All nine FRIENDS sessions were delivered to classes assigned to both the health-led and school-led conditions. To assess intervention fidelity, we recorded and independently rated 49 sessions (one from each class in the 28 schools delivering FRIENDS). All specified core tasks and home activities were delivered in the 24 health-led sessions. In the school-led sessions, 15 of 25 (60%) delivered all core tasks and the home activity, eight (32%) delivered all except the home activity, and two (8%) did not deliver one core task and the home activity. Session attendance was not recorded although average school absence rates were very low (4·2% in the health-led group and 4·4% in the school-led group).

We collected primary outcome data at 12 months from 1257 (92%) of the children who completed baseline assessments (449 [92%] in the health-led group, 436 [92%] in school-led group, and 372 [93%] in the usual school provision). We recorded a significant difference in adjusted mean RCADS at 12 months for health-led FRIENDS compared with school-led FRIENDS (interaction coefficient −3·91, 95% CI −6·48 to −1·35; p=0·0004) and usual school provision (−2·66, −5·22 to −0·09, p=0·043). The 95% CIs include our predefined clinically important difference of 3·6 points on the RCADS. Analysis of the RCADS subscales showed difference in generalised and social anxiety, but not depression (table 2).

Analysis of other secondary outcomes and parent and teacher completed measures identified no differences between treatment groups at 12 months (table 3).

As specified in the protocol, we did separate subgroup analysis in children with the highest 10% of baseline RCADS scores (high anxiety ≥49) and the remaining 90% (low anxiety ≤48, table 4).

We recorded significant within-group reductions for the high-risk group at 12 months but no effects between groups. For the low-risk group, we noted between-group differences in mean RCADS at 12 months (table 4). Adjusted mean differences showed an effect for health-led FRIENDS versus school-led FRIENDS (adjusted difference −3·78, 95% CI −6·16 to −1·40; p=0·003) and health-led FRIENDS versus usual school provision (−3·13, −5·61 to −0·65; p=0·015). This effect relates to a reduction in the health-led FRIENDS group on the social and generalised anxiety subscales (table 2).
In the low anxiety group, the standardised effect size of health-led FRIENDS compared with usual school provision (Cohen’s $d=0.22$, 95% CI 0.38–0.07) was small. The economic evaluation will be published separately.

Discussion
This is the first large pragmatic randomised trial comparing a universally provided classroom-based cognitive behaviour therapy anxiety prevention programme led by health and school staff with usual school provision. When we transferred health-led FRIENDS to everyday settings, the programme was more effective in the reduction of child-reported symptoms of anxiety than was school-led FRIENDS or usual school provision (panel 2). Although intervention leaders received the same initial training, our data suggest that a manualised programme might result in different outcomes depending on who delivers it. Although training teachers to deliver mental health programmes offers a potentially convenient low-cost sustainable option, our results show that this approach is not as effective as delivery by health professionals.

Further exploration identified two potentially important differences between health and school leaders. First, although treatment fidelity was high, the programme was more effective in the reduction of child-reported symptoms of anxiety than was school-led FRIENDS or usual school provision (panel 2). Although intervention leaders received the same initial training, our data suggest that a manualised programme might result in different outcomes depending on who delivers it. Although training teachers to deliver mental health programmes offers a potentially convenient low-cost sustainable option, our results show that this approach is not as effective as delivery by health professionals.

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The effect was also specific to child report because although parent and teacher ratings reduced over time, these changes were not significant. This finding might suggest that although children are feeling better, adults have not noticed any improvement in functioning or reduction in distress. However, only 42% of parents returned baseline questionnaires and as such these findings might not be representative of the total cohort. Similarly, the teacher assessment was global and might be insensitive to specific changes in anxiety symptoms. Alternatively, the absence of a parent and teacher effect might show the difficulty of assessment of changes in internal emotional symptoms and cognitions that are not directly observable.

The results of our study are consistent with those of reviews in which the intervention leader affected programme effectiveness, and with other implementation trials in which teacher-led FRIENDS was not effective. Our findings differ to those of the study by Barrett and Turner, which directly compared teacher and psychologist delivery of FRIENDS, although their study lacked statistical power. Overall, our study supports the growing evidence base for the use of the FRIENDS programme as an effective school-based anxiety prevention programme.

The intervention that we assessed was based on cognitive behaviour therapy, which is typically used to treat mental health disorders. However, for school-based universal emotional health programmes, most children will be healthy and will not need treatment. Furthermore, provision of individual treatment in a classroom context would not be appropriate. Therefore, it is important to emphasise that we did not provide treatment but used the cognitive behaviour therapy framework to help children to develop emotional, cognitive, and behavioural skills. Provision of life skills development within the school context fits with the growing recognition that schools are not just concerned with the development of academic skills, but also have an important role in the enhancement of emotional development in children.

Our study has many strengths. We used a manualised anxiety prevention programme that has been shown to be effective in schools and has been piloted in UK schools. Recruitment and retention were high, absenteeism was low, and programme fidelity was good. In terms of limitations, we relied on self-report measures and did not undertake any diagnostic interviews. Although anxiety symptoms were reduced in the health-led FRIENDS group, whether this reduction was indicative of changes in diagnostic status or impairment is unclear. Second, although our regional cohort included a representative sample of UK schools on several key dimensions, our sample of UK schools on several key dimensions, our cohort was less socially disadvantaged and had more white British participants than did the average UK state school. Therefore, whether similar results would be obtained with a more disadvantaged or ethnically diverse population is unclear. Similarly, although our schools were well matched across trial groups and our exploratory qualitative analysis identified few differences in school culture and ethos, these differences could have affected outcomes. Third, although the health-led and school-led FRIENDS facilitators had the same initial training and treatment fidelity was good, we did not directly assess how the intervention was delivered. Therefore, the differences between these groups might be indicative of differences in leader enthusiasm, confidence, and ability to engage and motivate students, which are skills that could be developed in school staff with additional training and supervision. Finally, although we reported symptoms at 12 months, we were unable to establish whether the improvements reported would be sustained over time. This point is especially important for prevention programmes in which the full extent of the preventive effects might take several years to emerge.

In summary, our results are encouraging and show that anxiety prevention programmes delivered in schools to children aged 9–10 years do reduce anxiety symptoms at 12 months. The finding that children with low symptoms benefited from the programme supports a universal approach, which also fits well with school timetables and organisational structures. However, our data suggest that the same programme can result in different effects depending on who delivers it. Further research is needed to explore the longer term effect, and the potential mediators and moderators of anxiety prevention programmes, to assess cost-effectiveness, and to establish whether anxiety prevention programmes are effective with more diverse and disadvantaged groups.

Contributors
PS, GT, RP, HD, RA, and NS conceived and designed the study. ES managed the trial and supervised data collection. GT undertook the statistical analysis. All authors had access to all study data and participated in interpretation of the findings, contributed core ideas and were involved in critically revising the paper for important intellectual content. All authors read and approved the final manuscript. PS was principal investigator and will act as guarantor for the paper.

Declaration of interests
We declare no competing interests.

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References


