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Abstract

Background: The Thinking Healthy Programme (THP) is recommended to treat perinatal depression in resourcelimited settings, but scale-up is hampered by a paucity of community health workers. THP was adapted for peerdelivery (THPP) and evaluated in two randomized controlled trials in India and Pakistan. Our aim was to estimate the effectiveness of THPP on maternal outcomes across these two settings, and evaluate effect-modification by country and other pre-defined covariates. Methods: Participants were pregnant women aged≥18 years with depression (Patient Health Questionnaire (PHQ-9) score≥10), randomized to THPP plus enhanced usual care (EUC) or EUC-only. Primary outcomes were symptom severity and remission (PHQ-9 score<5) 6 months post-childbirth. Secondary outcomes included further measures of depression, disability and social support at 3 and 6 months post-childbirth. Results: Among 850 women (280 India; 570 Pakistan), 704 (83%) attended 6-month follow-up. Participants in the intervention arm had lower symptom severity (PHQ-9 score adjusted mean difference -0.78 (95% confidence interval -1.47,-0.09)) and higher odds of remission (adjusted odds ratio 1.35 (1.02,1.78)) versus EUC-only. There was a greater intervention effect on remission among women with short chronicity of depression, and those primiparous. There were beneficial intervention effects across multiple secondary outcomes. Limitations: The trials were not powered to assess effect-modifications. 10-20% of participants were missing outcome data. Conclusions: This pooled analysis demonstrates the effectiveness, acceptability and feasibility of THPP, which can be scaled-up within a stepped-care approach by engaging with the existing health care systems and the communities to address the treatment gap for perinatal depression in resource-limited settings.

Keywords	Pregnancy; community health workers; depression; patient health questionnaire; Pakistan; India
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14 November 2019

Dear Dr Brambilla

We thank you and the reviewer for the consideration of our manuscript entitled "Effectiveness of the Thinking Healthy Programme for perinatal depression delivered through peers: pooled analysis of two randomized controlled trials in India and Pakistan", and for your helpful feedback.

We are pleased to submit our revised manuscript with tracked changes, along with a point-bypoint response to the feedback.

We thank you for your continued consideration of our manuscript and we look forward to hearing from you.

Yours sincerely

Fiona Vanobberghen





POINT BY POINT RESPONSES TO REVIEWERS

Comments from the editors and reviewers:

-Reviewer 1

- Since THP/ THPP is the main intervention in this study, the authors should describe more (in details) about this type of intervention in the Introduction or summarize a protocol in a Table. Response: We have added further details of THPP to the methods (pages 2-3 of the revised manuscript) including citing the complete description in the published trial protocol paper and main trial papers (line 53) and in the intervention development paper (line 63). Manuals are available on request, as indicated (lines 63-64). We took this opportunity to restructure the first part of the methods section to improve readability, separating out the design/setting, interventions and outcomes.

- In the Discussion, the authors should compare THPP to other standard psychological intervention in perinatal depression (e.g. CBT, IPT, etc) and may propose the mechanism by which THPP can help in decreasing chronicity and increasing remission of depression.

Response: We have added discussion of other psychological interventions to the discussion (page 11, lines 274-279). The mechanism of THPP was assessed through a pooled mediation analysis of the two trials, which found that the key mechanism was through increase in behavioral activation and social support (Singla et al., 2019). We have added this reference to the manuscript (line 276) where we mention behavioural activation.

Singla, D.R., MacKinnon, D.P., Fuhr, D.C., Sikander, S., Rahman, A., Patel, V., 2019. Multiple mediation analysis of the peer-delivered Thinking Healthy Programme for perinatal depression: findings from two parallel, randomised controlled trials. Br J Psychiatry 1–8. https://doi.org/10.1192/bjp.2019.184

HIGHLIGHTS

- In this pooled analysis of two randomized controlled trials, we found consistent benefits of the Thinking Healthy Programme adapted for delivery by Peers (THPP) on maternal outcomes
- THPP resulted in 35% higher odds of remission at 6 months post-childbirth compared to enhanced usual care only
- There was evidence of beneficial intervention effects across a number of secondary outcomes, and in repeated measures analyses, demonstrating feasibility and acceptability of this peerdelivered intervention, and external validity of our results

ABSTRACT

Background: The Thinking Healthy Programme (THP) is recommended to treat perinatal depression in resource-limited settings, but scale-up is hampered by a paucity of community health workers. THP was adapted for peer-delivery (THPP) and evaluated in two randomized controlled trials in India and Pakistan. Our aim was to estimate the effectiveness of THPP on maternal outcomes across these two settings, and evaluate effect-modification by country and other pre-defined covariates. Methods: Participants were pregnant women aged≥18 years with depression (Patient Health Questionnaire (PHQ-9) score≥10), randomized to THPP plus enhanced usual care (EUC) or EUC-only. Primary outcomes were symptom severity and remission (PHQ-9 score<5) 6 months post-childbirth. Secondary outcomes included further measures of depression, disability and social support at 3 and 6 months post-childbirth.

Results: Among 850 women (280 India; 570 Pakistan), 704 (83%) attended 6-month follow-up. Participants in the intervention arm had lower symptom severity (PHQ-9 score adjusted mean difference -0.78 (95% confidence interval -1.47,-0.09)) and higher odds of remission (adjusted odds ratio 1.35 (1.02,1.78)) versus EUC-only. There was a greater intervention effect on remission among women with short chronicity of depression, and those primiparous. There were beneficial intervention effects across multiple secondary outcomes.

Limitations: The trials were not powered to assess effect-modifications. 10-20% of participants were missing outcome data.

Conclusions: This pooled analysis demonstrates the effectiveness, acceptability and feasibility of THPP, which can be scaled-up within a stepped-care approach by engaging with the existing health care systems and the communities to address the treatment gap for perinatal depression in resource-limited settings.

Key words: Pregnancy; community health workers; depression; patient health questionnaire;

Pakistan; India

Trial registrations: ClinicalTrials.gov: NCT02104232, NCT02111915

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1 Effectiveness of the Thinking Healthy Programme for perinatal depression delivered through peers:

2 pooled analysis of two randomized controlled trials in India and Pakistan

3

4 INTRODUCTION

Depression is the main cause of disability globally, and its prevalence is increasing (Friedrich, 2017).
More than one in ten women experience a major depressive episode in the vulnerable perinatal
period (pregnancy or the year following delivery) in high income countries (Fisher et al., 2012;
Hendrick et al., 1998), with prevalences in low- and middle-income countries (LMIC) of up to 20%
(Fisher et al., 2012; Woody et al., 2017). Perinatal depression is associated with adverse maternal
outcomes including suicidal ideation (Gelaye et al., 2016), pregnancy and birth complications, and
poorer mental, motor and emotional development of the infant (Field et al., 2006).

12

13 Psychological therapies like cognitive behavior therapy or behavioral activation are recommended 14 for the treatment of mild and moderate perinatal depression (Guille et al., 2013). Due to the paucity 15 of mental health professionals in LMIC (Kakuma et al., 2011), some of these interventions are 16 modified, and provided as low-intensity psychological interventions delivered by trained para-17 professionals to depressed mothers in need (Rahman et al., 2013). One such low-intensity 18 psychological intervention is the Thinking Healthy Programme (THP), developed by Rahman and 19 colleagues (Rahman, 2007) and recommended by the World Health Organization (WHO) for the 20 treatment of perinatal depression in LMIC (World Health Organization, 2015). THP is delivered by 21 community healthcare workers (CHW) and has shown to be highly effective, halving the risk of 22 perinatal depression and significantly improving infant health outcomes in a study in Pakistan 23 (Rahman et al., 2008). However, the public health impact of THP has been hampered by the high 24 occupational load of CHWs (Haq et al., 2008), therefore impairing the scale up of this programme 25 (Jaskiewicz and Tulenko, 2012). To address this challenge, the National Institute of Mental Health 26 (NIMH), through the Collaborative Hubs for International Research on Mental Health (CHIRMH) 27 initiative, commissioned two studies in India and Pakistan to adapt the Thinking Healthy Programme

for delivery by trained peers, namely local women with no prior experience of healthcare delivery (Atif et al., 2017; D. Singla et al., 2014). Randomized controlled trials were conducted in two distinct settings, in urban/peri-urban Goa, India and rural Rawalpindi, Pakistan (Sikander et al., 2015), to evaluate the effectiveness and cost-effectiveness of the Thinking Healthy Programme Peer-delivered (THPP) compared to enhanced usual care (EUC). There were some differences in the trial designs (outlined below), but both trials found moderate effects of THPP on depression symptom severity and remission at 3 months post-childbirth (Fuhr et al., 2019; Sikander et al., 2019).

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Our aim was to estimate the overall effectiveness of THPP on maternal outcomes, and evaluate effect modifications by country and other pre-defined covariates. The increased power from the pooled analysis enables us to look for consistency of results across the two settings, to assess the external validity of the intervention effect, and hence the ability to generalize findings regarding acceptability and feasibility of this peer-delivered intervention to other settings.

41

42 METHODS

43 Trial settings, designs, and participants

44 Data were pooled from the Goa, India trial (hereafter, THPP-India) and the Rawalpindi, Pakistan trial 45 (hereafter, THPP-Pakistan). The trial settings are diverse, with higher rates of poverty (60% versus 46 4%), lower proportions of women educated (65% versus 84%), higher fertility rates (3.8 versus 1.8) 47 and larger households (6.2 versus 4.2 persons per household) in rural Rawalpindi compared to urban 48 Goa (D. Singla et al., 2014). The trial designs were similar (Table 1): both trials enrolled pregnant 49 women aged \geq 18 years with moderate to severe depression defined by scoring \geq 10 on the nine-item 50 Patient Health Questionnaire (PHQ-9). THPP-India was run through healthcare facilities with women 51 individually-randomized, while THPP-Pakistan was conducted in a community setting with woman 52 randomized in village clusters to avoid contamination. The trials have been described in full 53 elsewhere (Fuhr et al., 2019; Sikander et al., 2019, 2015).

55	Trial	interventions

- 56 <u>Participants were randomized to</u>, and compared THPP plus EUC to or EUC-only. The
- 57 interventions THPP consisted of 6-14 sessions over the prenatal period to six months post-childbirth,
- 58 covering behavioral activation, active listening, collaboration with the family, guided discovery and
- 59 homework (Atif et al., 2017). In India, THPP was delivered as 6-14 individual sessions, and in Pakistan,
- 60 THPP was delivered as ten individual and four group sessions. Peers received classroom and field
- 61 training, and received regular supervisions by the trainers. Emphasis was on behavior activation
- 62 strategies to enable the delivery by peers with no previous experience of delivering healthcare.
- 63 <u>Further details of THPP have been published previously (Atif et al., 2017) and ;</u> manuals are available
- 64 <u>from the authors on request.</u> THPP-India was run through healthcare facilities with women
- 65 individually-randomized, while THPP-Pakistan was conducted in a community setting with woman
- 66 randomized in village clusters to avoid contamination.
- 67

68 Trial outcomes

- 69 In both trials, the primary outcomes were symptom severity (PHQ-9 score) and remission (PHQ-9
- score <5) at 6 months post-childbirth. The trials have been described in full elsewhere (Fuhr et al.,

71 2019; Sikander et al., 2019, 2015). In this analysis, we included the primary and secondary maternal

- 72 outcomes.
- 73

74 Statistical methods

- 75 Baseline characteristics of the two trial populations were described, and a multivariable logistic
- 76 regression model was used to assess factors independently associated with country. Process
- indicators were summarized by country, including peer characteristics and attendance to therapy
- 78 sessions among women in the intervention arm.
- 79
- 80 Outcome data were analyzed using logistic and linear regression models to estimate odds ratios and
- 81 mean differences, respectively (with 95% confidence intervals). Generalized estimating equations

82 (GEEs) with an exchangeable correlation structure were used to account for the village-level 83 clustering in THPP-Pakistan (Sikander et al., 2019), with individuals in THPP-India acting as their own 84 clusters. Models were adjusted for all variables included in the main trials' analyses, because they 85 were: pre-specified in the analysis plan, stratification factors in the randomization, unbalanced 86 between groups at baseline, or associated with missing outcome data at 6 months. We also included 87 country. The following baseline variables were adjusted for as fixed effects: country, recruitment site, 88 residence (rural/urban), union council, symptom severity, treatment expectations, education, 89 chronicity of depression, and time between screening and birth. Participants with missing values 90 were omitted from the models. 91 92 For the primary outcomes, the following variables were assessed as a priori effect modifiers, by 93 fitting interactions between the intervention and the covariate: country, age, chronicity of 94 depression, baseline symptom severity, treatment expectations, and parity. We also assessed effect 95 modification by country for the secondary outcomes. Repeated measures analyses were performed, 96 combining the 3 and 6 months results, with assessment of group by time interactions. All models 97 were adjusted for the covariates listed above. 98 99 Sensitivity analyses were performed for the primary outcomes, using GEEs with individuals grouped 100 as one cluster, and mixed effects models. No adjustments were made for multiple testing; results 101 were interpreted based on the strength of evidence of effect size and consistency of results across outcomes. Analyses were conducted in Stata (StataCorp, 2015), following intention-to-treat 102 103 principles. Data are available on request (Fiona Vanobberghen et al., 2018; F Vanobberghen et al., 104 2018). 105 **Ethical considerations** 106

107 Ethical approval for the THPP-India trial was obtained from the Institutional Review Boards (IRBs) at

108 the London School of Hygiene and Tropical Medicine (LSHTM), Sangath (the trial-implementing

institution in India), and the Indian Council of Medical Research. Ethical approval for the THPPPakistan trial was obtained from the IRBs at the University of Liverpool, LSHTM, and the Human
Development Research Foundation (the trial-implementing institution in Pakistan). Participants in
both trials provided written informed consent (or witnessed informed consent/audio-recordings for
illiterate participants).

114

115 **RESULTS**

116 Overall, 280 women were enrolled in THPP-India and 570 in THPP-Pakistan (Table 2). After adjusting 117 for confounders (Appendix Table A1), THPP-Pakistan participants were less likely than those in THPP-118 India to work (6% versus 15%, with these low rates reflecting the national populations where the 119 majority of women are housewives) (WHO Country Office Pakistan, 2013; World Bank, 2013), had 120 received more education, had higher expectations of the usefulness of counseling, had higher 121 baseline symptom severity, had lower social support scores, were more likely to have had a previous 122 miscarriage or stillbirth, and were less likely to report domestic violence. Data on chronicity of 123 depression at baseline were missing for 30% of THPP-Pakistan and no THPP-India participants. There 124 were some differences between the women who did and did not have chronicity data in THPP-125 Pakistan: those with missing data were somewhat less likely to work, had higher treatment 126 expectations, had lower symptom severity, were more likely to be multiparous, were more likely to 127 have had a previous non-live birth, and were less likely to report domestic violence. Data were 128 missing on the time between screening and birth of the child for 15% of THPP-Pakistan and 4% of THPP-India participants, due to women being lost to follow-up after screening. Depression was more 129 130 chronic among THPP-Pakistan than THPP-India participants and time between screening and the 131 birth of the child was shorter.

132

Overall, 26 and 66 peers were trained and delivered at least one session in THPP-India and THPPPakistan, respectively, with corresponding mean ages of 38 and 30 years and mean years of
education completed 12 years in both trials (Table 3). No peers were lost during the THPP-India trial,

whereas 23 (35%) were lost during the THPP-Pakistan trial. Attendance to supervisions was 67% and
88% in THPP-India and THPP-Pakistan, respectively. Quality was assessed differently in the two trials.
In THPP-India, 18 items were assessed on the Therapy Quality scale of 0-2 (higher better) (D. R. Singla
et al., 2014); the overall score across 72 sessions rated by independent raters was 1.49 (standard
deviation 0.33). In THPP-Pakistan, assessments were done by independent raters for each peer in
three sessions at each of three time-points using an 18-item competency checklist; the overall
average was 84%. These results indicate average to good therapy quality.

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The mean number of sessions attended by participants in the intervention arm was slightly higher in THPP-Pakistan compared to THPP-India (10.9 versus 9.8); this was driven by higher attendance in the postnatal period (Table 3). In THPP-India, treatment completion was defined as attending at least six sessions with at least one session in each of four phases (prenatal, and 1-2, 3-4 and 5-6 months postnatal); 99/138 (72%) women completed treatment. In THPP-Pakistan, treatment completion was defined as attending at least ten sessions; 201/258 (78%) women completed treatment.

150

151 Average symptom severity decreased considerably over time in both groups (Appendix Figure A1). 152 There was evidence of a beneficial effect of the intervention on both primary outcomes at 6 months, 153 with participants in the intervention group having lower symptom severity (adjusted mean difference 154 -0.78, 95% confidence interval -1.47,-0.09) and higher odds of remission (adjusted odds ratio 1.35, 155 95% confidence interval 1.02,1.78) compared with those in the control group (Table 4, Figures 1A 156 and 1B). There was a trend towards stronger intervention effects in THPP-India compared to THPP-157 Pakistan for remission, but the difference was not statistically significant (p=0.18, with p=0.77 for 158 symptom severity). Results were robust to sensitivity analyses (Appendix Table A2).

159

After adjusting for confounders including country, there was evidence of a greater intervention effect on remission at 6 months among women with shorter chronicity of depression at baseline, and those primiparous (p=0.03 for both interactions, Figure 1B and Appendix Table A3). Similar trends were

163 observed for symptom severity with chronicity and parity, although the p values were large (p=0.49 164 and 0.30, respectively; Figure 1A and Appendix Table A4). For both outcomes, there were non-165 significant trends towards greater intervention effects among younger women (p=0.14 and 0.27 for 166 remission and symptom severity, respectively). There was no consistent evidence of effect-167 modification by baseline symptom severity or treatment expectations. 168 169 After adjusting for confounders including intervention group, the following factors were associated 170 with higher symptom severity and lower odds of remission at six months: country, higher baseline 171 symptom severity, and lower level of education (Appendix Table A5). 172 173 There was consistently strong evidence of beneficial intervention effects across a number of 174 secondary outcomes (Table 4). For symptom severity and WHO-DAS score, there were slightly 175 stronger intervention effects at 3 months than at 6 months. For remission and MSPSS score, results 176 were broadly similar at 3 and 6 months. For the composite outcomes of recovery and response at 177 both 3 and 6 months, there was consistent and strong evidence of a benefit of the intervention. 178 There was no evidence of an intervention effect on number of days unable to work in the last month. 179 There was no evidence of effect modification by country for any of the secondary outcomes. 180 181 There was no statistical evidence of group by time interactions, and assuming a constant intervention 182 effect at 3 and 6 months (Appendix Table A6), there were strong evidence of intervention effects on symptom severity, remission, WHO-DAS score and MSPSS score (3 and 6 months combined), in line 183 184 with the results of the individual trials (Fuhr et al., 2019; Sikander et al., 2019). There was no 185 evidence of an intervention effect on number of days unable to work in the last month. 186 DISCUSSION 187 188 We present results of a pooled analysis of two of the largest trials evaluating a psychological

189 intervention for perinatal depression in LMIC delivered by peers. With the increased power from this

pooled analysis, we found small (Rahman et al., 2013) but important benefits of THPP across a range
of maternal depression, disability and social support outcomes, with 35% higher odds of remission at
6 months post-childbirth compared to EUC-only. These findings open the possibilities of peerdelivered THPP to be implemented in other settings where there are limited numbers of mental
health professionals to tackle the treatment gap for perinatal depression.

195

196 Historically, intervention studies of depression in LMIC have generally had small sample sizes, with 197 different recruitment mechanisms of study participants and using varying methodological designs 198 and outcome measures (Chowdhary et al., 2014; Rahman et al., 2013). Our studies represent a new 199 generation of complementary trials of psychological therapies nested in the community using 200 innovative delivery mechanisms based on lay care providers and peer supervisors. THPP-India was 201 conducted in a peri-urban setting in Goa while THPP-Pakistan was embedded in rural communities of 202 Rawalpindi. These contextual and socio-economic differences were reflected in the trial populations 203 and delivery agents (peers), with the THPP-Pakistan participants having more severe and more 204 chronic depression at baseline compared to those in the THPP-India trial. THPP-Pakistan participants 205 reported on average more years of education than those in THPP-India, contrary to expected (D. 206 Singla et al., 2014) and compared to lower nationwide literacy rates of 45-50% (Population Census 207 Organization, 1998; World Bank, 2013). There may have been over-reporting of education among the 208 participants in Pakistan due to social desirability. Participants in THPP-Pakistan also had poorer 209 depression outcomes at 6 months compared to those in THPP-India, after adjustment for 210 confounders including baseline severity and chronicity. Slightly different models of incentivization 211 were used across the two trials: peers in urban Goa received financial incentives since this was 212 identified as an important motivator, while peers in rural Rawalpindi were volunteers, devoting their 213 time to THPP for altruistic motives (Fuhr et al., 2019; Sikander et al., 2019; D. Singla et al., 2014). This 214 may have been one reason for a greater turnover of peers during the trial in Pakistan (35% compared 215 to none in India), though attendance to supervision was excellent (88%, compared to 67% in India). 216 Furthermore, most peers in Pakistan who left were able to identify their own replacements,

217 indicating that there was a pool of women interested in this work. The health systems in which peers 218 were trained to operate THPP were also distinct. Peers in Rawalpindi worked alongside government 219 lady health workers delivering care in the community, while peers in Goa worked more 220 independently and within a tiered public healthcare system (D. Singla et al., 2014). There were some 221 differences in the design of the intervention between the two trials (Sikander et al., 2015), although 222 the mean number of sessions attended by intervention recipients was similar. Despite these contextual differences, the impact of THPP by country was very similar, demonstrating the external 223 224 validity of the intervention. In both countries, peers were trained to competently deliver THPP, with 225 high proportions of women in the intervention arms completing their treatment. This indicates that 226 delivery by peers was not seen as stigmatizing, and demonstrates a high degree of acceptability of a 227 psychological therapy delivered by peers in community settings to mothers with depression in both 228 contexts. Further, the intervention was shown to offer an appreciable improvement in health at low 229 cost in THPP-Pakistan (incremental cost-effectiveness ratio US\$ 15.50 over the whole trial period) 230 and even cost saving in THPP-India (-US\$ 93.53) (Fuhr et al., 2019; Sikander et al., 2019), despite 231 monetary incentivisation for the peers contributing to 12% of the cost in the latter trial (D. Singla et 232 al., 2014).

233

234 While the benefits of THPP were consistent for remission across three and six months post-childbirth, 235 we observed stronger intervention effects at three compared to six months for symptom severity 236 (PHQ-9 score) and disability (WHO-DAS). Similar effects were found in the individual trials (Fuhr et 237 al., 2019; Sikander et al., 2019). This may be explained by the intervention design which puts 238 emphasis on the first three months after child-birth by front-loading sessions antenatally and soon 239 after childbirth, and/or by the natural remission rates of untreated depression over time (Austin et 240 al., 2008; Posternak and Miller, 2001; Rojas et al., 2007; Tandon et al., 2018; Whiteford et al., 2013). 241 The benefits at three months remain important to reduce the duration of the depression episode at 242 such a critical time-point in childcare. Considering effect modification of the intervention, we 243 observed greater benefits of THPP among women with shorter chronicity of depression at baseline

244 and those primiparous. THPP may be best adopted as a first-step psychological intervention to be 245 used in a stepped care system for maternal depression in LMIC, most suitable for lower-risk groups. 246 The need for a stepped collaborative care model is highlighted by the fact that nearly half of women 247 in the intervention arm across both countries did not achieve remission by 6 months. Future research 248 should explore if peers are able to sign-post higher risk women, and refer them to primary or 249 specialist care where they may receive additional interventions. However, we did not observe a 250 difference in the benefits of THPP by baseline severity, in contrast to previous studies of THP (Patel et 251 al., 2017).

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253 There are some limitations of this study. The control group received enhanced usual care in a well-254 resourced setting, thus the intervention effects might be greater in a health system where no such 255 care is provided (Fuhr et al., 2019). The individual trials were not powered for assessing effect 256 modifications; while the pooling of the data from the two trials provides more power we cannot rule 257 out having missed some effect modifications, including those by country. Some baseline covariates 258 were missing, most notably chronicity of depression in 30% of THPP-Pakistan participants. Outcome 259 data were missing for approximately 10% of THPP-India participants, and 20% of THPP-Pakistan 260 participants, although we adjusted for factors associated with missingness which yielded similar 261 results to multiple imputation analyses in the individuals trials (Fuhr et al., 2019; Sikander et al., 262 2019).

263

THPP-India and THPP-Pakistan were pragmatic trials with high internal validity delivered in community settings to mothers recruited from routine health-care settings. By aligning THPP with routine clinical practice and working alongside CHWs and clinicians, we were able to evaluate reallife treatment effects in two diverse South Asian contexts. This pooled analysis demonstrates the effectiveness, acceptability and feasibility of the THP intervention delivered by peers. This is important because although the theoretical model was the same in the two settings, there were contextual differences in implementation. The fact that the intervention effects were similar in the

two trials demonstrates the external validity of the intervention, and hence potential forgeneralizability to other settings and populations.

273

274 <u>THP uses cognitive behaviour therapy techniques and was designed to be delivered by lady health</u>

275 workers (Atif et al., 2017). THPP represents a further adaptation, with focus on behaviour activation

276 strategies to enable successful delivery by peers (Atif et al., 2017; Singla et al., 2019). Other

277 interventions for perinatal depression include educational sessions led by healthcare professionals,

278 or interpersonal therapy with problem-solving methods through group activities (Rahman et al.,

279 2013) but to our knowledge none have been adapted for delivery by non-healthcare professionals.

Although the benefits of THPP were relatively small (Rahman et al., 2013), it has proven to be cost-

281 effective (Fuhr et al., 2019; Sikander et al., 2019). These results therefore open avenues for

282 collaborative care, where peers can be the "missing" community agent or case manager to increase

awareness and provide first-line interventions, and work alongside specialists in primary, secondary

and tertiary care. Further, the delivery through peers provides opportunities for women in the

285 community to develop skills and improve their chances of employability. To make this happen, there

is a need in global mental health to advocate for accreditation of evidence-based and cost-effective

287 low-intensity psychological interventions like THPP. We conclude that THPP is suitable for

implementation as part of a stepped-care approach by engaging with the existing health-care

systems and the communities to treat perinatal depression in other contexts.

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504 FIGURE LEGENDS

505

506 Figure 1. A) Remission (PHQ-9 score <5) and B) symptom severity (PHQ-9 score) at 6 months by

507 potential effect modifiers.

508 PHQ=Patient Health Questionnaire. Results are from linear or logistic generalized estimating equation models, adjusted for 509 country, recruitment site, residence, union council, baseline symptom severity, treatment expectations, education,

- 510 chronicity of depression, and time between screening and birth. The sizes of the squares indicating the point estimates are
- 511 proportional to the number of participants in each category. These results are described further in Table 4 and Appendix
- 512 Tables A3 and A4.
- 513
- 514

TABLES

Table 1. Summary of the trial designs.

	THPP-India	THPP-Pakistan			
Location	Goa	Rawalpindi			
Study setting	Antenatal clinics in two hospitals and three primary health centers	Community setting in rural sub-district			
Participants	Pregnant women aged ≥18 years with moderate to severe depression as defined by PHQ-9 score ≥10				
Intervention	THPP delivered as 6-14 individual sessions, each lasting 30-45 minutes, in four phases (prenatal, up to two months post childbirth, 3-4 months after childbirth, and 5-6 months after childbirth), plus enhanced usual care	THPP delivered as ten individual and four group sessions, each lasting 30-45 minutes, from the third trimester of pregnancy to six months after childbirth, plus enhanced usual care			
Control	Enhanced usual care: Standard care from the gynecologist plus (1) patients and gynecologists were informed of participants screening results; (2) gynecologists were given the adapted mental health Gap Action Programme (mhGAP) treatment guidelines for perinatal depression; (3) participants were provided with an information sheet on health-care during pregnancy and beyond	Enhanced usual care: Standard care from LHWs plus (1) all participants and LHWs were informed of participants screening results; (2) doctors and midwives at the primary healthcare centers were given the adapted mental health Gap Action Programme (mhGAP) treatment guidelines for perinatal depression; (3) participants were provided with an information sheet on healthcare during pregnancy and beyond			
Peers	Women with children, a similar socio- demographic background as participants, and good communication skills. Recruited from the local community through word-of-mouth, particularly through key informants in women's self- help groups and CHWs. Financial incentives were paid based on sessions delivered, up to a maximum of Rs. 5900 (approximately US\$80) per trial participant, plus a monthly honorarium of Rs. 100 (approximately US\$1) to cover phone call costs	Women with children, a similar socio- demographic background as participants, and good communication skills. Identified through LHWs and elders, with recruitment and placement through primary health-care centers. Peers received a travel allowance for supervision meetings and a phone card/top-up to be able to stay in touch with their respective supervisors in case of support needed. No remunerations were paid for delivering sessions			
Randomization	Individually-randomized (1:1 allocation ratio), stratified by place of residence	Cluster-randomized (1:1 allocation ratio; 40 village clusters), stratified by 11 union			
Blinding Primary outcomes Secondary outcomes ^a	(rural/urban) councils Outcome assessors were blinded to participants' treatment allocation Symptom severity (PHQ-9 score) and remission (PHQ-9 <5) at 6 months post-childbirth Symptom severity (PHQ-9 score) at 3 months post-childbirth Remission (PHQ-9 score <5) at 3 months post-childbirth Recovery (PHQ-9 score <5 at both 3 and 6 months post-childbirth) Response I (PHQ-9 score <10 at both 3 and 6 months post-childbirth) Response II (≥50% reduction from baseline PHQ-9 score at both 3 and 6 months post- childbirth) WHO-DAS score, at 3 and 6 months post-childbirth Number of days unable to work in the last month, at 3 and 6 months post-childbirth				
Recruitment period and follow up Number of	October 2014-June 2017 280	October 2014-March 2017 570			

- The trial designs have been described previously in detail (Sikander et al., 2015). CHW=community health workers. LHW=
- Lady Health Worker. THPP=Thinking Healthy Programme Peer-delivered. PHQ=Patient Health Questionnaire.
- 518 519 520 MSPSS=Multidimensional Scale of Perceived Social Support. WHO-DAS=World Health Organization Disability Assessment
- 521 Schedule. ^aMaternal secondary outcomes only.

523 **Table 2. Baseline characteristics by trial.**

	THPP-India	THPP-Pakistan	Total
Number enrolled	280	570	850
Age, years (mean (SD; range))	25 (4.6; 18-41)	27 (4.8; 18-45)	26 (4.8; 18-45)
Level of education (n (%))			
No formal education	34 (12%)	107 (19%)	141 (17%)
Up to primary	120 (43%)	39 (7%)	159 (19%)
Up to secondary	90 (32%)	333 (58%)	423 (50%)
Beyond secondary	36 (13%)	91 (16%)	127 (15%)
Occupation (n (%))			
Does not work	237 (85%)	533 (94%)	770 (91%)
Works	43 (15%)	37 (6%)	80 (9%)
Chronicity of depression, weeks (n (%)) ^a			
<12	173 (62%)	73 (18%)	246 (36%)
≥12	107 (38%)	326 (82%)	433 (64%)
Symptom severity (PHQ-9 score) (median	12 (11,15)	14 (12,17)	14 (11,17)
(IQR))			
Symptom severity category (n (%))			
Moderate (PHQ-9 score 10-14)	197 (70%)	312 (55%)	509 (60%)
Moderately severe (15-19)	67 (24%)	187 (33%)	254 (30%)
Severe (20-27)	16 (6%)	71 (12%)	87 (10%)
MSPSS score (mean (SD))	5 (1.1)	4 (1.4)	4 (1.4)
Participant's expectation of usefulness of	counseling (n (%))	b	
Not useful	1 (0%)	4 (1%)	5 (1%)
A little	55 (20%)	26 (5%)	81 (10%)
somewhat useful	54 (19%)	114 (20%)	168 (20%)
Moderately useful	58 (21%)	246 (43%)	304 (36%)
Very useful	112 (40%)	178 (31%)	290 (34%)
Parity (n (%))			
Primiparous	119 (43%)	102 (18%)	221 (26%)
Multiparous	161 (57%)	468 (82%)	629 (74%)
Previous miscarriage or still birth (n (%))			
None	261 (93%)	377 (66%)	638 (75%)
One/more	19 (7%)	193 (34%)	212 (25%)
Any domestic violence in last three month	ıs (n (%))⁰		
No	243 (87%)	486 (87%)	729 (87%)
Yes	37 (13%)	71 (13%)	108 (13%)
Time between screening and birth of	4 (1.6)	3 (1.2)	3 (1.5)
child, months (mean (SD)) ^d			

524 Percentages are of non-missing values. PHQ=Patient Health Questionnaire. MSPSS=Multidimensional Scale of Perceived

525 Social Support. SD=standard deviation. ^aMissing for 171 THPP-Pakistan participants. ^bMissing for 2 THPP-Pakistan

participants. ^cMissing for 13 THPP-Pakistan participants. ^dMissing for 11 THPP-India and 85 THPP-Pakistan participants.
 527

529 Table 3. Process indicators.

	THPP-India	THPP-Pakistan
Peers		
Peers trained and delivered at least one session	26	66
(number)		
Age of peers, years (mean, SD)	38 (7.5)	30 (5.5)
Years of education completed by peers (mean,	12 (2.7)	12 (2.5)
SD)		
Peers lost during the trial (number, %)	0	23 (35%)
Attendance to supervisions (%)	67%	88%
Participants in the intervention arm		
Number ^a	138	258
Sessions attended (mean, SD) ^b	9.8 (4.3)	10.9 (3.9)
Prenatal sessions attended (mean, SD) ^c	4.5 (1.9)	3.7 (1.7)
Postnatal sessions attended (mean, SD) ^d	5.4 (2.9)	7.3 (2.7)
Attended at least 6 sessions	113 (82%)	230 (89%)
Attended at least 10 sessions	87 (63%)	201 (78%)
Completed treatment ^e	99 (72%)	201 (78%)

530 SD=standard deviation. ^aExcluding 2 women in THPP-India and 25 in THPP-Pakistan who were discontinued as per protocol

before month 3 due to child death, still birth or abortion. ^bOf possible 14 sessions in each trial. ^cOf possible 6 sessions in

532 THPP-India and 5 in THPP-Pakistan. ^dOf possible 8 sessions in THPP-India and 9 in THPP-Pakistan. ^eDefined in THPP-India as 533 attended at least 6 sessions, with at least 1 session in each of four phases (prenatal, and 1-2, 3-4 and 5-6 months postnatal);

defined in THPP-Pakistan as attended at least 10 sessions.

536 Table 4. Primary and secondary outcomes.

	Number of participants ^a		Mean (SE) or number (%)ª		Intervention effect (adjusted mean difference or odds ratio: 95% CI) ^b	P value
	Control	Interven- tion	Control	Interven- tion		
Primary outcomes						
Symptom severity (PI	HQ-9 scor	e) at 6 mont	hs			
Overall	355	349	6.0 (0.3)	5.1 (0.3)	-0.78 (-1.47,-0.09)	0.03
India	129	122	4.5 (0.4)	3.5 (0.4)	-0.95 (-2.31,0.41)	
Pakistan	226	227	6.8 (0.4)	6.0 (0.4)	-0.72 (-1.52,0.08)	
Remission (PHQ-9 sco	ore <5) at	6 months	. ,			
Overall	355	349	178 (50%)	201 (58%)	1.35 (1.02,1.78)	0.04
India	129	122	77 (60%)	89 (73%)	1.86 (1.08.3.20)	
Pakistan	226	227	101 (45%)	112 (49%)	1.20 (0.87.1.66)	
Secondary outcomes	;				(, , , , , ,	
Continuous variables	5					
Symptom severity	333	346	7.1 (0.4)	5.5 (0.3)	-1.84 (-2.431.25)	<0.001
(PHO-9 score) at 3			/ = (,	010 (010)		
months						
WHO-DAS score						
3 months	332	346	17.0 (1.0)	14.2 (0.9)	-3 17 (-5 73 -0 61) ^c	0 02°
6 months	354	349	160(10)	13.6 (0.9)	-2 11 (-4 77 0 55)	0.02
Number of days	034	017	10.0 (1.0)	10.0 (0.7)	2.11(1.77,0.00)	0.12
unable to work in						
last month						
3 months	333	346	17(03)	16(03)	-0.24 (-0.86 0.39)	0.46
6 months	357 357	340	1.7(0.3)	1.6 (0.3)	0.24 (0.00, 0.37)	0.40
MSDSS score	334	547	1.0 (0.2)	1.0 (0.3)	0.02 (-0.31,0.33)	0.75
2 months	222	245	47(01)	10(01)	0 17 (0 02 0 21)	0.01
6 months	252	240	4.7 (0.1)	4.7 (0.1)	0.17(0.03, 0.31)	0.01
Categorical variables	554	347	4.7 (0.1)	5.0 (0.1)	0.27 (0.11,0.43)	0.001
	222	246	155 (170/)	106 (510/)	1 24 (1 12 1 60)	0.001
Q<5) at 2 months	333	340	133 (4770)	100 (54%)	1.34 (1.13,1.00)	0.001
	217	225	100 (22%)	125 (120/)	1 60 (1 21 2 10)	0.001
Recovery (PHQ-9<5	317	325	100 (32%)	135 (42%)	1.80 (1.21,2.10)	0.001
at Doth 3 and 0						
monuns)	217	225	100 (500/)	224 (60%)	1 57 (1 16 0 10)	0.002
Response I (PHQ-	31/	325	199 (24%)	224 (09%)	1.57 (1.10,2.10)	0.003
9 < 10 at both 3 and						
o montns)	047	005	4 4 4 4 5 9 ()	400 (500/)		.0.001
Response II (≥50%	31/	325	144 (45%)	190 (58%)	1.69 (1.27,2.24)	<0.001
reduction from						
baseline PHQ-9 at						
both 3 and 6						
months)						

537 Health Questionnaire. MSPSS=Multidimensional Scale of Perceived Social Support. WHO-DAS

538 Organization Disability Assessment Schedule. SE=standard error. CI=confidence interval. ^aIndia and Pakistan trials combined

539 for the secondary outcomes. ^bLinear or logistic GEE models, adjusted for country, recruitment site, residence, union council,

540 baseline symptom severity, treatment expectations, education, chronicity of depression, and time between screening and

541 birth (see methods). Baseline treatment expectations missing for one woman in Pakistan control group, therefore models

542 are based on one fewer woman than the numbers indicated. Country-level intervention effects are based on models with

543 interaction between group and country. ^cUsing independent correlation matrix due to convergence problems when using

544 exchangeable correlation structure.





Declaration of interest

Declarations of interest: none.

Contributors

DCF, BW, AL, HAW, AR and VP designed the THPP-India trial. EA, ADS, AJ, PK, and RK were responsible for intervention implementation and data gathering instruments in the THPP-India trial. BW, AL, and VP were responsible for THPP-India trial conduct. ED'S designed and managed the database for the THPP-India trial. SS, IA, DCF, HAW, VP and AR designed the THPP-Pakistan trial. IA, NA, AB, TB, SB, RL, MS and SZ were responsible for intervention implementation and data gathering instruments in the THPP-Pakistan trial. SS, AZ, and AR were responsible for THPP-Pakistan trial conduct. AZ designed and managed the database for the THPP-Pakistan trial. FV and HAW performed the statistical analyses. VP and AR provided oversight. FV and HAW drafted the manuscript. All authors read and approved the manuscript.

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pooled analysis of two randomized controlled trials in India and Pakistan

SUPPLEMENT

Table A1. Factors	associated	with country.
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Variable	Adjusted odds ratio (95% CI)	P value
	for Pakistan versus India ^a	
Age, per 10 years	0.98 (0.50,1.91)	0.95
Level of education		<0.001
None	1 (reference)	
Primary	0.09 (0.04,0.24)	
Secondary	1.60 (0.73,3.50)	
Higher secondary/above	2.46 (0.96,6.28)	
Occupation		0.003
Does not work	1 (reference)	
Works	0.28 (0.12,0.65)	
Chronicity of depression, weeks		<0.001
<12	0.14 (0.08,0.25)	
≥12	1 (reference)	
Symptom severity (PHQ-9 score)		0.003
10-14	1 (reference)	
15-19	2.41 (1.32,4.40)	
≥20	3.73 (1.29,10.81)	
MSPSS score, per unit	0.40 (0.31,0.51)	<0.001
Participant's expectation of usefulness of counselling		<0.001
Not/a little useful	1 (reference)	
Somewhat useful	3.30 (1.14,9.55)	
Moderately useful	11.17 (3.97,31.43)	
Very useful	5.07 (1.89,13.60)	
Parity		0.56
Primiparous	1 (reference)	
Multiparous	1.22 (0.63,2.36)	
Previous miscarriage or still birth		<0.001
None	1 (reference)	
One or more	9.43 (4.05,21.95)	
Domestic violence		0.01
No	1 (reference)	
Yes	0.34 (0.15,0.79)	
Time between screening and birth of child, per month	0.54 (0.43.0.66)	< 0.001

PHQ=Patient Health Questionnaire. MSPSS=Multidimensional Scale of Perceived Social Support. Cl=confidence interval. ^aLogistic regression model. N=589 (women with any missing data were excluded). Broadly similar results were obtained from models excluding chronicity of depression and time between screening and birth (which had the highest missingness, particularly in Pakistan).

Table A2. Sensitivity analyses for the primary outcomes.

	Number of clusters	Intervention effect (adjusted mean difference or odds ratio; 95% CI) ^a	P value	P value for effect modification by
				country
Symptom severity (PHQ-9 score) at 6 months				
Primary analysis as reported in Table 4 (GEEs with	291	-0.78 (-1.47,-0.09)	0.03	0.77
individuals in India acting as their own clusters)				
GEEs with individuals in India grouped as one cluster	41	-0.82 (-1.64,-0.01) ^b	0.05 ^b	0.81 ^b
Random effects models with individuals in India acting as	291	-0.82 (-1.64,-0.01)	0.05	0.81
their own clusters				
Random effects models with individuals in India grouped as	41	-0.82 (-1.64,-0.01)	0.05	0.81
one cluster				
Remission (PHQ-9 score <5) at 6 months				
Primary analysis as reported in Table 4 (GEEs with	291	1.35 (1.02,1.78)	0.04	0.18
individuals in India acting as their own clusters)				
GEEs with individuals in India grouped as one cluster	41	1.39 (1.02,1.91) ^b	0.04 ^b	0.20 ^b
Random effects models with individuals in India acting as	291	1.39 (0.02,1.91)	0.04	0.20
their own clusters				
Random effects models with individuals in India grouped as	41	1.39 (1.02,1.91)	0.04	0.20
one cluster				

^aLinear or logistic GEE models as described in the methods section, adjusted for country, recruitment site, residence, union council, baseline symptom severity, treatment expectations, education, chronicity of depression, and time between screening and birth (see methods). ^bUsing independent correlation matrix due to convergence problems when using exchangeable correlation structure.

	Number o	f participants	ts Number with remission (%)		Intervention effect (adjusted odds ratio; 95% CI)ª	P value for effect modification
	Control	Intervention	Control	Intervention		
Age, years						0.18
18-24	135	132	91 (69%)	74 (55%)	1.86 (1.08,3.20)	
≥25	220	217	110 (51%)	104 (47%)	1.20 (0.87,1.66)	
Chronicity of depression, weeks						0.03
<12	105	103	73 (71%)	55 (52%)	2.54 (1.39,4.65)	
≥12	172	178	88 (49%)	79 (46%)	1.10 (0.73,1.65)	
Missing ^b	78	68	40 (59%)	44 (56%)	-	
Baseline symptom severity (PHQ-9	score)					0.49
10-14	223	209	135 (65%)	124 (56%)	1.48 (1.01,2.17)	
≥15	132	140	66 (47%)	54 (41%)	1.17 (0.73,1.90)	
Treatment expectations ^c						0.85
None/little/somewhat	114	96	56 (58%)	58 (51%)	1.29 (0.73,2.26)	
Moderate/very useful	241	252	144 (57%)	120 (50%)	1.37 (0.98,1.93)	
Parity						0.03
Primiparous	91	95	72 (76%)	52 (57%)	2.59 (1.35,4,93)	
Multiparous	264	254	129 (51%)	126 (48%)	1.10 (0.80,1.52)	

Table A3. Remission (PHQ-9 score <5) at 6 months by potential effect modifiers.

PHQ=Patient Health Questionnaire. CI=confidence interval. ^aLogistic GEE models, adjusted for country, recruitment site, residence, union council, baseline symptom severity, treatment expectations, education, chronicity of depression, and time between screening and birth (see methods). Baseline treatment expectations missing for one woman in Pakistan control group, therefore models are based on one fewer women than indicated by the numbers indicated. These results are illustrated in Figure 1B. ^bModel includes missing chronicity category, but interaction effects are presented only among those with non missing chronicity. ^cMissing for one woman in Pakistan control group, who is omitted.

	Number of participants		Mean symptom severity		Intervention effect (adjusted	P value for effect
			(PHQ-9 score; SE)		mean difference; 95% CI) ^a	modification
	Control	Intervention	Control	Intervention		
Age, years						0.27
18-24	135	132	5.9 (0.5)	4.4 (0.5)	-1.38 (-2.66,-0.10)	
≥25	220	217	6.0 (0.4)	5.5 (0.4)	-0.44 (-1.35,0.47)	
Chronicity of depression, weeks						0.49
<12	105	103	5.1 (0.5)	3.7 (0.5)	-1.67 (-3.16,-0.18)	
≥12	172	178	6.9 (0.5)	5.8 (0.4)	-0.99 (-2.02,0.03)	
Missing ^b	78	68	4.9 (0.6)	5.6 (0.7)	-	
Baseline symptom severity (PHQ-9	score)					0.80
10-14	223	209	5.1 (0.4)	4.4 (0.4)	-0.69 (-1.65,0.27)	
≥15	132	140	7.4 (0.6)	6.2 (0.5)	-0.91 (-2.14,0.31)	
Treatment expectations ^c						0.35
None/little/somewhat	114	96	6.3 (0.6)	4.5 (0.5)	-1.37 (-2.80,0.05)	
Moderate/very useful	241	252	5.8 (0.4)	5.3 (0.4)	-0.54 (-1.39,0.30)	
Parity						0.29
Primiparous	91	95	4.8 (0.5)	3.3 (0.5)	-1.51 (-3.05,0.04)	
Multiparous	264	254	6.4 (0.4)	5.8 (0.4)	-0.51 (-1.32,0.29)	

PHQ=Patient Health Questionnaire. SE=standard error. CI=confidence interval. ^aLinear GEE models, adjusted for country, recruitment site, residence, union council, baseline symptom severity, treatment expectations, education, chronicity of depression, and time between screening and birth (see methods). Baseline treatment expectations missing for one woman in Pakistan control group, therefore models are based on one fewer women than indicated by the numbers indicated. These results are illustrated in Figure 1A. ^bModel includes missing chronicity category, but interaction effects are presented only among those with non missing chronicity. ^cMissing for one woman in Pakistan control group, who is omitted.

	Symptom severity (PHQ-9 score)		Remission (PHQ-9 score <5) at		
	at 6 months		6 months		
	Adjusted mean difference (95% CI) ^a	P value	Adjusted odds ratio (95% CI)ª	P value	
Randomised group		0.03		0.04	
Control	0 (reference)		1 (reference)		
Intervention	-0.78 (-1.47,-0.09)		1.35 (1.02,1.78)		
Country		<0.001		0.001	
India	0 (reference)		1 (reference)		
Pakistan	2.81 (1.23,4.39)		0.34 (0.18,0.64)		
Recruitment site		0.84		0.81	
N/A (Pakistan) or Goa Medical College Hospital	0 (reference)		1 (reference)		
Asilo	0.32 (-1.15,1.79)		1.10 (0.62,1.96)		
Primary health centres	-0.46 (-3.93,3.02)		1.62 (0.37,7.20)		
Residence		0.67		0.85	
N/A (Pakistan) or rural	0 (reference)		1 (reference)		
Urban	-0.38 (-2.09,1.33)		1.07 (0.54,2.09)		
Union council		0.002		0.03	
N/A (India) or Guf	0 (reference)		1 (reference)		
Kallar	-0.61 (-2.22,0.99)		1.44 (0.76,2.75)		
Kanoha	0.06 (-1.70,1.83)		1.47 (0.71,3.02)		
Sakot	-1.67 (-3.81,0.46)		1.55 (0.66,3.65)		
Choa	-1.33 (-3.23,0.58)		1.00 (1.00,1.00)		
Smoot	-2.60 (-4.66,-0.53)		1.87 (0.86,4.09)		
Bewek	1.68 (0.15,3.21)		3.14 (1.32,7.47)		
Darkali	-0.92 (-2.38,0.54)		0.57 (0.29,1.11)		
Bishandot	0.49 (-1.19,2.17)		1.24 (0.69,2.24)		
Sagri	-0.84 (-2.39,0.71)		1.31 (0.67,2.56)		
Nallah	-2.57 (-4.75,-0.39)		1.52 (0.81,2.86)		
Symptom severity (PHQ-9		0.004		0.02	
score)					
10-14	0 (reference)		1 (reference)		
15-19	1.56 (0.61,2.51)		0.59 (0.41,0.85)		
≥20	1.39 (-0.10,2.87)		0.71 (0.40,1.26)		
Participant's expectation		0.78		0.99	
of usefulness of					
	0 (mafamamaa)		(/ /		
Somewhat useful	-0.22 (-1.01,1.30)		0.99 (0.53,1.84)		
	-0.19 (-1.70,1.33)		0.95 (0.52,1.72)		
Education status	-0.02 (-2.12,0.00)	0.000	1.01 (0.50,1.82)	0.12	
No formal education	0 (reference)	0.009	1 (roforonco)	0.12	
Up to primary	-0.37 (-1.03, 1.07)		1.12 (0.04,1.70)		
Beyond secondary	-0.47 (-1.00,0.71)		1.43 (U.71,2.27)		
Chronicity of depression	2.30 (-3.70,-0.00)	0.12	1.75 (1.07,3.35)	0.07	
weeks		0.13		0.07	

Table A5. Factors associated with the primary outcomes.

<12	-0.45 (-1.52,0.63)		1.06 (0.70,1.62)	
≥12	0 (reference)		1 (reference)	
Missing	-1.23 (-2.44,-0.01)		1.74 (1.09,2.79)	
Time between screening	-0.16 (-0.46,0.14)	0.30	1.08 (0.96,1.22)	0.19
and birth of child, per				
month				

PHQ=Patient Health Questionnaire. MSPSS=Multidimensional Scale of Perceived Social Support. CI=confidence interval. N/A=not applicable. ^aLinear or logistic GEE models as described in the methods section, adjusted for all variables shown in the table.

Table A6. Analy	vsis of repeate	ed measures fo	or the primary	v and secondary	voutcomes.
Table AV. Allal	ysis of repeat	u measures n	or the primar	y and secondar	y outcomes.

	P value for group by time interaction	Overall adjusted mean difference or odds ratio (95% CI)	P value for overall intervention effect
Primary outcomes			
Symptom severity (PHQ-9 score)	0.17	-1.30 (-1.86,-0.74)	<0.001
Remission (PHQ-9 score <5)	0.98	1.39 (1.13,1.72)	0.002
Secondary outcomes			
WHO DAS score	0.81	-2.56 (-4.31,-0.81)	0.004
Number of days unable to work in last month	0.82	-0.12 (-0.55,0.32)	0.59
MSPSS score	0.37	0.25 (0.11,0.38)	<0.001

PHQ=Patient Health Questionnaire. WHO-DAS=World Health Organization Disability Assessment Schedule.

MSPSS=Multidimensional Scale of Perceived Social Support. CI=confidence interval. Linear or logistic GEE models, adjusted for visit month, country, recruitment site, residence, union council, baseline symptom severity, treatment expectations, education, chronicity of depression, and time between screening and birth (see methods).



Figure A1. Symptom severity (PHQ-9 score) over time.

The central line shows the median score, the length of the box is the interquartile range, the lines are the most extreme values within 1.5 times the interquartile range, and points outside this range are shown individually.