**TITLE PAGE**

**Evaluating feasibility and acceptability of a group WHO trans-diagnostic intervention for women with common mental disorders in rural Pakistan: a cluster randomised controlled feasibility trial**

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**Abstract**

**Aims:** The aim of this feasibility trial was to evaluate the feasibility and acceptability of the locally adapted Group PM+ intervention for women in the conflict affected settings in Swat, Pakistan.

**Methods:**

This mixed-methods study incorporated a quantitative component consisting of a two arm cluster randomised controlled feasibility trial, and qualitative evaluation of the acceptability of the Group PM+ to a range of stakeholder groups. For the quantitative component, on average from each of the twenty Lady Health Workers (LHWs) catchment area (20 clusters), 6 women were screened and recruited for the trial with score of > 2 on the General Health Questionnaire (GHQ-12) and score of > 16 on the WHO Disability Assessment Schedule (WHODAS). These LHW clusters were randomised on a 1:1 allocation ratio using a computer based software through a simple randomisation method to the Group PM+ intervention or Enhanced Usual Care. The Group PM+ intervention consisted of 5 weekly sessions of two hours duration delivered by local non-specialist females under supervision. The primary outcome was individual psychological distress, measured by levels of anxiety and depression on the Hospital Anxiety and Depression Scale at 7th week after baseline. Secondary outcomes include symptoms of depression, Post-Traumatic Stress Disorder (PTSD), general psychological profile, levels of functioning and generalized psychological distress. Intervention acceptability was explored through in-depth interviews.

**Results:**

The results show that lay-helpers with no prior mental health experience can be trained to achieve the desired competency to successfully deliver the intervention in community settings under supervision. There was a good intervention uptake, with Group PM+ considered useful by participants, their families, and lay-helpers. The outcome evaluation, which was not based on a large enough study to identify statistically significant results, indicated statistically significant improvements in depression, anxiety, general psychological profile and functioning. The PTSD symptoms and depressive disorder scores showed a trend in favour of the intervention.

**Conclusion:**

This trial showed robust acceptance in the local settings with delivery by non-specialists under supervision by local trained females. The trial paves the way for further adaptation and exploration of the outcomes through larger-scale implementation and definitive Randomized Controlled Trials (RCTs) in the local settings.

**Trail registration:** Australian New Zealand Clinical Trials Registry: ACTRN12615000210572 (<https://www.anzctr.org.au/Trial/Registration/TrialReview.aspx?id=368082>). Registered 5th March 2015.

**Introduction**

Research on mental health problems[[1]](#footnote-1) following humanitarian emergencies[[2]](#footnote-2) has focused on post-traumatic stress disorder (PTSD) ([George et al., 2012](#_ENREF_15), [Inter-Agency Standing Committee, 2007](#_ENREF_19)). However, a broad range of mental health consequences are recognised ([World Health Organization, 2007](#_ENREF_44)). While the prevalence rates of mental health problems during and after humanitarian emergencies tend to be higher compared to other populations, there is considerable variation ([Inter-Agency Standing Committee, 2007](#_ENREF_19), [Steel et al., 2009](#_ENREF_39)). The average rates for depression and PTSD range between 15-20% ([Steel et al., 2009](#_ENREF_39)). Women are especially vulnerable to common mental disorders due to lack of resources, low education, limited social support and gender-based violence ([Allden et al., 1996](#_ENREF_2), [Bhui et al., 2003](#_ENREF_7), [Ellsberg et al., 2008](#_ENREF_14)). A recent survey (current study site) found 38% of women were psychologically distressed, and associated with previous exposure to conflict-related trauma and major stressors such as death/disability of a significant other, financial constraints, accommodation difficulties and interpersonal problems including marital issues ([Khan et al., 2015](#_ENREF_22)).

The evidence base regarding psychological treatments for mental health problems specifically related to trauma is mounting, however, the majority of this research has been conducted in high income settings or through specialized professionals ([Tol et al., 2011](#_ENREF_40), [Tol et al., 2013](#_ENREF_41)). In order to be scalable in humanitarian emergencies, it is important to establish whether such interventions remain effective when delivered by non-specialists under supervision. Such interventions has received significant attention as part of the global mental health research agenda ([Kakuma et al., 2011](#_ENREF_20)). These interventions should be brief and simple so that they can be delivered by people in the community ([Rahman et al., 2008](#_ENREF_34)). Additionally, interventions should address a range of outcomes (trans-diagnostic), including physical functioning and common mental health and psychosocial problems.

The WHO, as part of its Mental Health Gap Action Programme (mhGAP), has developed guidance for humanitarian settings and is testing psychological interventions delivered by non-specialists, including Problem Management Plus (PM+) ([Dawson et al., 2015](#_ENREF_12), [Ventevogel et al., 2015](#_ENREF_42)). PM+ has been developed for adults affected by symptoms of common mental health problems including depression, anxiety, and those related to practical issues like unemployment and interpersonal relationship difficulties, within contexts following exposure to an emergency.

In order to fit different service, contexts and beneficiary groups, it will be important to have both individual and group versions of PM+. This will aid accessibility by a range of target groups. The group format draws on the same strategies as the individual version of PM+ but also has the benefits of peer interaction and the potential for group members to be therapeutic agents to each other ([Yalom & Leszcz, 2005](#_ENREF_46)). A Meta-analysis suggests that group psychotherapeutic interventions based upon cognitive behavioural approaches are as effective as individual interventions at 6-month follow-up ([Cuijpers et al., 2008](#_ENREF_11)). Due to the ability to reach larger numbers of people, the group format offers a potentially scalable approach that may also be a cost-effective means of intervention.

The aim of this study was to evaluate the feasibility and acceptability of the Group PM+ intervention in rural Swat which has recently been exposed to conflict. A key feature was to explore the feasibility of non-specialist delivery of the intervention in collaboration with the Lady Health Workers (LHWs) programme[[3]](#footnote-3).

**Methods**

**Study design**

This mixed-methods study incorporates a two arm cluster randomised controlled feasibility trial, and qualitative evaluation of the acceptability of Group PM+ intervention to a range of stakeholders.

**Settings and participants**

Swat is a rural district in the northern part of the Khyber Pakhtunkhwa province, Pakistan. The majority of the population (86%) lives rural ([Pakistan Bureau of Statistics, 2016](#_ENREF_30)). Since 2007 Swat has witnessed widespread conflict/militancy. During the active conflict between the army and militants in 2009, nearly 2.5 million people of the district were internally displaced across Pakistan ([Bile & Hafeez, 2009](#_ENREF_8)). In July 2009, these displaced persons were asked to repatriate after the Government declared the area safe, although continued military presence and sporadic conflicts between militants and security agencies continues to create social unease ([Rome, 2010](#_ENREF_36)).

The current trial was conducted from 21st March until 12th August, 2015 in Qambar Union Council (UC), a rural conflict-affected area in Swat. Each UC has a Basic Health Unit (BHU) staffed by a physician, a Lady Health Visitor (primary healthcare worker stationed at a health unit and providing maternal and child health services), a vaccinator, a midwife and LHWs. LHWs are trained to provide mother and child healthcare and education to a catchment area of approximately 1,000 people or 150 homes, conducting monthly routine health visits ([Hafeez et al., 2011](#_ENREF_18)). Given their existing roles, LHWs provided a link between Group PM+ facilitators/delivery agents and women in the community, overcoming a significant access barrier.

Participants were females aged 18 years or above, referred for screening based on the judgment of their LHW that they were psychologically distressed. Participants who scored above 2 on the General Health Questionnaire (GHQ) and above 16 on the WHO Disability Assessment Schedule (WHO-DAS) were invited to participate in the trial by trained assessment team. Assessment team members were females from the local community. Those Women with imminent suicide risk, cognitive impairment (e.g., severe intellectual disability or dementia), or mental disorder (psychotic disorders, substance-dependence) were excluded.

As a cluster-randomized control trial the unit of randomization was the LHWs catchment area. Of the 24 LHWs in Qambar UC, 20 LHWs were eligible for randomization (Figure 1), which was carried out by an independent researcher. Based on 1:1 allocation ratio, 10 LHWs were randomized to the intervention and 10 to Enhanced Usual Care (EUC)[[4]](#footnote-4) arm. On average six women were recruited for the trial from each LHW catchment area. When randomized into the intervention arm, the LHW along with the lay-helper informed the participant of the schedule for five consecutive weekly Group PM+ sessions, with the first session scheduled not longer than two weeks after the pre-intervention assessment. The two lay-helpers carried out half of the intervention sessions each.

**Intervention**

**Group Problem Management Plus:** PM+ is a WHO psychological intervention developed on evidence of established problem solving, counselling and behavioural techniques ([Dawson et al., 2015](#_ENREF_12)). Group PM+ is an adaptation of the individual intervention. The multicomponent nature of PM+ allows for many symptoms of common mental health disorders to be addressed with the one intervention (i.e. trans-diagnostic). The manual has been translated into Urdu and Pashtu, and culturally adapted to the Swat setting. This study is the first testing of Group PM+ which comprises five weekly sessions each lasting two hours, inclusive of breaks. All LHWs in the intervention arm were provided a half-day training session by the lay-helpers regarding their roles and responsibilities in the trial. LHWs initially served to facilitate introduction of lay-helpers to community, encourage participant attendance, and provide a space in which to conduct sessions. LHWs were present when Group PM+ sessions were conducted, although they were not active facilitators of the intervention content.

Group PM+ was delivered by local female lay-helpers with 16 years of education (graduates) and with no formal training of or prior experience in mental health. Lay-helpers and participants were gender matched to enhance acceptability and ensure researcher safety ([Craig et al., 2000](#_ENREF_10)). The training and supervision of lay-helpers followed an apprenticeship model ([Murray et al., 2011](#_ENREF_29)) which involves moving the primary delivery of healthcare from specialists such as psychiatrists and psychologists to lay-helpers, and building skills through on-the-job training. Three non-specialist supervisors (HN, PA and AM) and 2 female lay-helpers received 6 days training by the Master Trainer (KD). Training covered knowledge of common mental disorders, basic counselling and group management skills, the Group PM+ intervention and self-care strategies. This was followed by four weeks of practice cases with weekly group supervision through Skype (2-3 hours duration) by the 3 supervisors. This group supervision model was continued throughout the trial, providing opportunities for peer learning and collective problem-solving. Before delivering Group PM+ as part of the trial, lay-helpers completed competency assessments (explained below under process evaluation). These were developed specifically for PM+ and involved role-plays that were scored on fidelity to the treatment, competency of intervention delivery and counselling skills.

**Enhanced Usual Care (EUC)**

In Swat, treatment-as-usual in primary healthcare centers (PHCs) to individuals with common mental disorders usually consists of (a) no treatment, (b) placebo-based care, as evidence-based mental health care is not currently available in PHCs, or (c) referral to the District Headquarter Hospital (DHQ) for specialized psychiatric care. For this study, care was enhanced in two ways: (a) LHWs received training in primary care referral pathways for treatment of common mental disorders; and (b) primary care physicians received the training in assessment and treatment of common mental disorders routinely taught by our partner, the WHO Collaborating Center in Rawalpindi, Pakistan. Both of these trainings were delivered by the Swat District Psychiatrist in separate half-day sessions. During the study any EUC participants with severe psychiatric disorders (e.g., psychosis) or problems (e.g., suicidality) that required immediate specialist treatment and follow-up were referred to the DHQ or BHU, depending upon their needs.

**Measures**

Assessments were conducted at baseline and 7 weeks after baseline assessment. The instruments have been adapted and validated for use in Pakistan. Blinding was ensured during assessments through LHW instructing participants not to disclose their allocation status. In case of disclosure the assessments were reschedule through the other assessor.

*Anxiety and Depression (primary outcome measure):* Psychological distress in terms of states of anxiety and depression were measured with the Hospital Anxiety and Depression Scale ([Mumford et al., 1991](#_ENREF_28), [Zigmond & Snaith, 1983](#_ENREF_47)), an established scale consisting of 14 items. The HADS contains two sub-scales: HADS-A (anxiety, 7 items and range 0-21) and HADS-D (depression, 7 items and range 0-21). Higher scores indicate more severe symptoms of anxiety and/or depression. The HADS has been translated into Urdu, and showed satisfactory reliability and validity ([Mumford et al., 1991](#_ENREF_28)).

*Psychological distress (screening instrument):* This was measured by the GHQ at baseline assessment only ([Goldberg & Williams, 1988](#_ENREF_16), [Minhas & Mubbashar, 1996](#_ENREF_25)). When used as a screening tool, the GHQ is usually scored bi-modally (i.e., 0-0-1-1), with scores ranging from 0-12. In a previous study in Pakistan, a cut-off of 3 or higher has been reported to indicate clinically significant psychological distress ([Minhas & Mubbashar, 1996](#_ENREF_25)).

*Functioning (screening instrument):* Disability was assessed using the 12-item interviewer-administered screener version of the WHO Disability Assessment Schedule (WHODAS 2.0) ([World Health Organization, 2010](#_ENREF_45)). WHODAS 2.0 covers six domains (cognition, mobility, self-care, getting along, life activities and participation), and assesses difficulties people have during the last 30 days. Difficulties are scored on a 5-point Likert scale as none, mild, moderate, severe, or extreme. WHODAS 2.0 has been validated cross-culturally and is valid as a screening and outcome measure, displaying good sensitivity to change ([Andrews et al., 2009](#_ENREF_3), [Sousa et al., 2010](#_ENREF_38)).

*Depressive disorder:* The Patient Health Questionnaire (PHQ-9) is a 9-item instrument measuring presence and severity of depressive disorder ([Kroenke et al., 2001](#_ENREF_23)). As a severity measure, the PHQ-9 score ranges from 0 to 27, with each of the 9 items scored from 0 (not at all) to 3 (nearly every day). The PHQ has been validated in Urdu ([Ahmer et al., 2007](#_ENREF_1)).

*Post-traumatic stress symptoms:* PTSD symptoms were measured using the PTSD Checklist for DSM-5 ([Weathers et al., 2013](#_ENREF_43)). This 20-item checklist corresponds with the 20 symptoms of the DSM-5 PTSD diagnosis. Items were rated on a 0-4 scale, with a total severity score of 80. The previous version of the PCL based on DSM IV PTSD symptoms, the PCL-C, has been used previously in Pakistan ([Khalily et al., 2012](#_ENREF_21)). The PCL-5 was adapted to ask for symptoms in the last week (rather than month) to enhance sensitivity to change.

*Psychological Outcome Profiles (PSYCHLOPS):* PSYCHLOPS consists of four questions ([Ashworth et al., 2004](#_ENREF_6)), covering three domains: problems (2 questions), function (1 question) and wellbeing (1 question). Responses are scored on an ordinal six-point scale producing a maximum score of 18 (6 points per domain). Lower scores indicate better psychological profile. PSYCHLOPS gave participants the opportunity to identify problems they wish to address and discuss during group session. Although PSYCHLOPS has not been validated in South Asian context, the tool has shown good convergent validity, satisfactory internal consistency with measures of psychological distress, and high sensitivity to change ([Ashworth et al., 2009](#_ENREF_5)).

Adverse life events: Potentially traumatic events, day to day life events, and demographic information was also collected to compare the two arms for major baseline differences ([Mollica et al., 1992](#_ENREF_27), [Rahman et al., 2003](#_ENREF_33)).

**Process evaluation**

The process evaluation involved mixed-methods process monitoring and semi-structured process evaluation interviews. Process monitoring involved analysis of supervision records and assessments of lay-helper competency. The lay-helpers were assessed by four independent assessors through application of a specially developed competency rating tool that evaluated a) counselling skills, and b) use of PM+ strategies with participants through direct observation of specially designed role plays. Competency was rated using a 5-point Likert scale ranging from 0 (not done) to 5 (excellent).

The qualitative evaluation explored questions relating to intervention acceptability through targeted semi-structured interviews with key informants following an established approach for applied mental health research in humanitarian settings ([Applied Mental Health Research Group, 2013](#_ENREF_4), [Green & Thorogood, 2014](#_ENREF_17)). Interviews followed a semi-structured topic guide developed for each category of key informant including intervention participants, lay-helpers, LHWs, family members of intervention participants and assessment team members. They were all interviewed through the semi structured topic guide on feasibility/acceptability of the trial procedures in the community. Interviews were documented verbatim in a written transcript for subsequent analysis.

**Data analysis**

Since this was a feasibility trial, no power calculations were carried out ([Leon et al., 2011](#_ENREF_24)). Feasibility testing of all the components of training, delivery and supervision requires the LHWs have a sufficient case-load. The sample size allowed each LHW to have 6 cases (one group) which is sufficient for a feasibility study.

We carried out intention-to-treat analysis to assess the intervention effect (participant were randomized and analyzed irrespective of whether or not they received the allocated intervention as long as they completed the end point assessment). A linear mixed model was used for the analysis of continuous outcomes which had treatment as a fixed effect, baseline measurement of the primary outcome as a covariate, and cluster as random effect. The treatment difference in least square means (treatment effect) between the two arms together with its 95% confidence interval was derived from the mixed model.

All qualitative data was analysed using thematic content analysis ([Applied Mental Health Research Group, 2013](#_ENREF_4), [Green & Thorogood, 2014](#_ENREF_17), [Ritchie et al., 1994](#_ENREF_35)) relevant to applied research. This involved researchers familiarizing themselves with interview transcripts, coding transcripts by themes and sub-themes, and finalizing coding through consensus. A document containing thematic categories was then reviewed alongside interview transcripts to identify quotes to illustrate key themes. All analysis was conducted in the original language, with only the quotes presented here translated into English.

Ethical approval was obtained from the local Ethics Review Board at the Lady Reading Hospital and the WHO Ethical Review Committee. Participants were only enrolled after providing voluntary written informed consent which included information on trial procedure, confidentiality, data management and reporting.

**Results**

A total of 134 participants referred by their respective LHWs were approached for screening. Fourteen participants declined participation before screening. The 120 participants who agreed to participate were screened on the two screening instruments after informed consent. After screening positive, 119 participants were invited for the trial. Of the 119, 59 participants in 10 LHW catchment areas were in the intervention arm, and 60 participants in 10 LHW catchment areas were in the control arm. Seven participants of the 119 randomized (6%) were lost to follow-up. See figure 1 for a detailed consort flow diagram. Table 1 shows the sample characteristics, and demonstrates the treatment arms were well balanced at baseline for demographic variables.

Table 2 summarizes results for the primary and secondary outcomes in the intervention and control arms on intention-to-treat analysis. On the HADS score (primary outcome measure), there was a difference in the scores of both arms with the intervention arm mean HADS score 4.73 less than the control group (95% CI: (-7.41,-2.06, p=0.001). This difference in scores was observed for the separate anxiety and depression scores, at 2.62 (95% CI: (-4.46,-0.77, p=0.008) and 2.48 (95% CI: (-4.00,-0.97, p=0.002) respectively. Similarly on the WHODAS (functionality), PSYCHLOPS, PTSD symptoms (PCL) and depressive disorder (PHQ) scores showed a trend in favor of intervention.

Although the study was not powered to detect clinically significant differences on outcome measures, the results indicated that PM+ had the potential to improve functioning, symptoms of anxiety and depression.

Process evaluation

Process monitoring demonstrated that participants showed good levels of intervention adherence with 36/59 (61%) attending all five sessions; and 14/59 (24%), 4/59 (6.8%), 3/59 (5%) and 1/59 completing four, three, two, and one session/s respectively. Initially, both lay-helpers achieved unsatisfactory results in their competency assessments. Additional targeted training and supervision (two days over Skype sessions and visits) was therefore provided to support in areas where their scores were low, alongside additional practice sessions in the field under intensive supervision. Following this the lay-helpers were re-assessed and deemed competent to deliver Group PM+.

Key objectives of the qualitative interviews were to explore: The acceptability of the intervention to key stakeholders, the feasibility of LHWs supporting integration of the intervention into their routine scope of work and identifying barriers and challenges to further scale-up. Interviews were conducted with two lay-helpers, five participants who completed the sessions, four family members of the completing participants, three LHWs, and two assessment team members. The number of interviews was predetermined and weighted for each category of respondent. Due to time constraints data collection did not reach saturation (no new information after data saturation).

Interviewees raised common themes which are summarised in Table 3. Overall lay-helpers, participants, and their families expressed satisfaction with Group PM+, identifying its ability to address problems relating to participants’ perceived psychological well-being and daily functioning. Furthermore, respondents felt that lay-helpers developed good rapport with the participants.

A barrier identified by participants and family members was the duration of sessions (2 hours). Furthermore, participants expectation of monetary incentives to attend sessions (in line with the routine practice of many international organization programmes), led to difficulties motivating participants to attend, and may have influenced lower levels of intervention attendance. Finally, confidentiality issues arose in sessions containing multiple participants from the same household. Consequently, many participants felt uncomfortable disclosing their problems to the group, instead seeking to discuss these with the lay-helper separately during breaks.

**Discussion**

This mixed-method study assessed the acceptability and feasibility of Group PM+ through a feasibility cluster randomized controlled trial. The results show that lay-helpers can be trained to achieve the desired competency to successfully deliver the intervention in community under supervision. Overall, intervention uptake was good, with Group PM+ considered useful by participants, their families, and lay-helpers. The outcome evaluation, which was not powered, nonetheless indicated relevant improvements in symptoms of depression, anxiety; general psychological profile and functioning. Barriers identified included confidentiality issues which suggest that sessions need to be comprised of women from different households. A recommendation is that the length of the session be reduced through shortening session breaks to minimise the time that females were away from household responsibilities.

A major strength of this study is that it has been conducted in a post-conflict area with adequate recruitment and low dropout rate. The intervention was found to be acceptable to the local community and LHWs. Furthermore, there was a high trial retention rate (94%) which further suggests community acceptability of the intervention ([Ehlers et al., 2005](#_ENREF_13), [Rachman et al., 2008](#_ENREF_31)). This could be due to the accessibility of the intervention in the community which reduced the potential stigma associated with seeking mental health support ([Rahman et al., 2013](#_ENREF_32)). Indeed, good retention was also found in a trial of group interpersonal psychotherapy in Uganda ([Bolton et al., 2003](#_ENREF_9)). The feasibility of procedures and acceptability of the intervention to the community, LHWs and health system show that larger scale definitive trials are possible in the setting.

LHWs successfully facilitated the introduction of lay-helpers to the community and invited participants to the sessions. They were ideal hosts for Group PM+ as they are trusted and respected in the communities, overcoming a barrier to accessing women in need. Rural Swat may be considered a highly conservative society and without the LHWs as gatekeepers, community members are unlikely to trust and accept the research procedures and new health interventions ([Mohmand & Gazdar, 2007](#_ENREF_26)).

The group format of the intervention had some inherent challenges; notably that women were reluctant to discuss their problems in the presence of family members. For the definitive trial this challenge will be addressed by separating family members into different PM+ groups. In addition, the long duration of each session for the definitive trial will be managed by either reducing the duration of breaks or taking them at the end of the session, according to the preference of each group.

The lay-helpers were recruited from the local area with the aim of ensuring acceptability and an ability to relate well to their participants. There is data showing that participants can have preference for delivery agents from within the communities over outside specialists, as they are easily accessible and reduce the stigma attached to help-seeking ([Singla et al., 2014](#_ENREF_37)). Furthermore, local selection is considered to enhance the potential for future intervention scale-up. The lay-helpers were graduates with no prior experience of mental health interventions, enabling access to a large pool of potential persons to take on this role. In addition, locally embedded lay-helpers could be considered more conscious of the local context, including awareness of cultural norms and sensitivity to potential security concerns.

The supervision structures are potentially scalable, with a small number of trained specialists in another centre/city providing group-supervision to many lay-helpers through video calls. Additionally, the group format of supervision enabled lay-helper peer support and supervision which was found to be important for managing difficult situations arising during field activities, including lay-helpers emotional wellbeing. Training of lay-helpers was followed by four weeks of practice cases with weekly group supervision, therefore to enhance transfer of skills, more emphasis on experiential components and role-plays should be considered.

In conclusion, the current study shows encouraging results regarding the feasibility and acceptability of the Group PM+ intervention delivered by lay-helpers. The intervention showed improvement in depression, anxiety and functioning, although the small sample size means the results need to be interpreted with caution, with a larger scale definitive trial required to demonstrate intervention effectiveness.

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**Conflicts of Interest**

None.

**Ethical Standards**

The authors assert that all procedures contributing to this work comply with the ethical standards of the relevant national and institutional committees on human experimentation and with the Helsinki Declaration of 1975, revised 2008.

**Authorship**

Research was conceptualized by AR & MvO. All authors have reviewed and approved the final manuscript for submission.

**Availability of data and material**

Raw data is available as additional supporting files.

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***Figure 1: Consort flow diagram of the feasibility trial***

**Enrolment**

No. of clusters assessed for eligibility n= 24

Excluded clusters n= 4

LHWs not fully inducted n= 2

LHWs not able to take part due to personal reasons n= 2

Randomized n= 20

**Allocation**

Analysis done n= 58

7th week post assessment n= 58 (97%)

Lost to follow up n= 2 (3%)

Clusters allocated to control n= 10

Average cluster size n=6

Participants approached n= 68

Refused before screening n= 8

Participants screened n= 60

Participants enrolled for the trial n= 60

**Analysis**

Analysis done n= 54

**Follow up**

7th week post assessment n= 54 (92%)

Refused n= 2 (3%)

Lost to follow up n=3 (5%)

Clusters allocated to intervention n= 10

Average cluster size n=6

Participants approached n= 66

Refused before screening n= 6

Participants screened n= 60

Participants negatively screened n= 1

Participants enrolled for the trial n= 59

Participants attended 4-5 sessions= 50/59 (85%)

Participants attended 2-3 sessions= 7/59 (12%)

Participants attended $\leq $1 session= 2/59 (3%)

**Table 1: Demographic variables between intervention and control group**

|  |  |  |
| --- | --- | --- |
| **Variables** | **Intervention group (N=59)****n (%)** | **Control group (N=60)****n (%)** |
| **Age** |  |  |
| 25 years or less | 21 (36) | 16 (27) |
| 26-35 years | 18 (31) | 19 (32) |
| 36-45 years | 18 (31) | 21 (35) |
| 46 years or more | 2 (3) | 4 (7) |
| **Marital status** |  |  |
| Never married | 7 (12) | 7 (12) |
| Currently married | 41 (70) | 47 (78) |
| Separated | 3 (5) | 3 (5) |
| Widowed | 8 (14) | 3 (5) |
| **Education** |  |  |
| None | 41 (70) | 41 (68) |
| Primary | 9 (15) | 10 (17) |
| Secondary | 6 (10) | 6 (10) |
| 10 years or above | 3 (5) | 3 (5) |
| **Occupation** |  |  |
| Paid work | 10 (17) | 16 (27) |
| House keeping  | 48 (81) | 41 (68) |
| **Respondent husband’s occupation** |  |  |
| Paid work | 19 (43) | 13 (28) |
| Self-employed | 18 (41) | 22 (47) |
| Unemployed | 1 (2) | 2 (4) |
| **Financially empowered**  | 27 (46) | 26 (43) |
| **Nuclear family structure** | 16 (27) | 16 (27) |
| **Family size** |  |  |
| ≥2 children younger than 7 years old | 17 (29) | 22 (37) |

**Table 2: Comparison of outcomes between control and intervention groups on intention to treat analysis**

| **Primary and secondary outcomes** | **Time** | **N** | **Intervention group****Mean (SD)** | **N** | **Control****group****Mean (SD)** | **Linear Mixed Model** |
| --- | --- | --- | --- | --- | --- | --- |
| **Difference in least squares mean (95%CI)** | **p-value** |
| Hospital Anxiety & Depression Rating Scale (Depression and Anxiety combined) | Baseline   | 59 | 20.15(7.13) | 60 | 24.15(7.58) | -4.65(-7.35,-1.95) | 0.0009 |
| One Week post intervention | 54 | 15.70(9.04) | 58 | 23.21(8.32) |
| Hospital Anxiety Rating Scale (Anxiety score) | Baseline   | 59 | 10.81(3.68) | 60 | 13.13(3.94) | -2.62(-4.37,-0.86) | 0.0039 |
| One Week post intervention | 54 | 7.59(4.66) | 58 | 11.52(4.59) |
| Hospital Depression Rating Scale (Depression score) | Baseline  | 59 | 9.34(4.38) | 60 | 11.02(4.32) | -2.48(-4.00,-0.96) | 0.0016 |
| One Week post intervention | 54 | 8.11(5.13) | 58 | 11.69(4.42) |
| WHO DAS (Functionality) | Baseline | 59 | 30.24(6.62) | 60 | 31.92(7.20) | -5.37(-8.97,-1.76) | 0.0040 |
| One Week post intervention | 54 | 24.44(8.90) | 58 | 30.86(8.64) |
| PSYCHLOPS | Baseline | 55 | 14.69(3.31) | 60 | 15.63(3.39) | -4.49(-6.41,-2.58) | <0.0001 |
| One Week post intervention | 54 | 9.22(4.97) | 57 | 14.21(3.98) |
| PCL (PTSD symptoms) | Baseline  | 59 | 28.47(15.80) | 60 | 36.53(16.73) | -2.79(-9.51,3.94) | 0.4128 |
| One Week post intervention | 54 | 17.65(15.59) | 58 | 24.02(16.26) |
| PHQ (Generalized distress) | Baseline  | 59 | 11.42(5.71) | 60 | 13.23(6.92) | -1.06(-3.59,1.48) | 0.4112 |
| One Week post intervention | 54 | 10.06(6.45) | 58 | 12.05(6.09) |

**Table 3: Qualitative interview results**

|  |  |  |
| --- | --- | --- |
| **Objective** | **Theme** | **Quote (source)** |
| Acceptability to participants | Participants learnt useful skills to manage their problems | *This was a very good experience. I am satisfied. We learned some techniques here that we didn’t knew before. We discussed our problem in groups and tried to solve them. (Client)**This was easy for me to implement the skills because these were solving my problems and my tension and the heaviness of my head improved, so I practised them regularly. (Client)**I had very good relation with my helper and I liked her because she came for us, we learned a lot that changed our lives, I trusted her and told her all my problems, our tensions were reduced because of her, she taught us such skills that we didn’t knew, and those skills solved a lot of our problems, it changed our behaviours. (Client)* |
| Acceptability to families | Recognizable improvement in condition | *I think this is a very good program because the participant from our house improved a lot. (Client family)**My family realized that these sessions are regarding health so I was allowed to attend the sessions and they trusted the lay-helper. (Client)* |
| Acceptability to lay-helpers | Lay-helpers learnt the skills | *At the start it was difficult because I have no idea of the work. But then it all went well. I not only managed peoples’ problems and learnt the skills but also learned to manage my own routine problems. (Lay-helper)**Very friendly, they trusted us, they frankly shared their problems with us. If they could not share any problem during the session in front of other participants they spoke to us during break time. (Lay helper)**The participant realized that it is working and their small problems started resolving, this motivated them to continue. (Lay helper)* |
| Feasible for LHWs to act as intervention facilitators | LHWs respected and trusted in the community | *I had a very good experience with the program; as all of the participants gathered in my health house, collectively listened everyone problems and then discussed the solutions. I even learnt new skills which improved my own health as well. (LHW)**I am officially working in this catchment area, so all these people know me and trust me. That’s the reason they send their women to my house. (LHW)* |
| Barriers and challenges | Confidentiality issues | *If in any session there were two or three participants from the same house that made it difficult to let the participants share their problems. (Lay helper)**I was not able to share my problems because of my family member in the same group. (Client)* |
| Monetary expectations  | *Participants wanted some monetary incentives and when it was not provided they lost interest and they were not punctual. (Lay helper)**Everyone believed that there would be some monetary incentives for the participants, but there was none. (Client)*  |
| Long session duration | *There was plenty of household chores to be done in the house, so at times the family got annoyed, the session duration was too long. (Client)**The sessions were too long and families have reservations, as the participants have to do work back home. (Lay helper)* |

1. Mental health problems: Mental health issues like anxiety, stress, Post Traumatic Stress Disorder (PTSD) and depression. [↑](#footnote-ref-1)
2. Humanitarian emergencies: Armed conflicts and natural disasters (including food crises) with large segments of populations at acute risk of death, immense suffering and/or losing their dignity. [↑](#footnote-ref-2)
3. Lady Health Workers (LHWs) programme was started in 1994 to provide health education, promote healthy behaviors, supply family planning methods and provide basic curative services in the communities. [↑](#footnote-ref-3)
4. Enhanced Usual Care (EUC): Usual care was enhanced by training primary care staff and ensuring referral pathways for common mental health problems. [↑](#footnote-ref-4)