



# Young adults as community mental health workers in humanitarian settings

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## Main Information

**Primary registry identifying number**

LBCTR2023015206

**Protocol number**

R34MH121558

**MOH registration number**

**Study registered at the country of origin**

Yes

**Study registered at the country of origin: Specify**

**Type of registration**

Prospective

**Type of registration: Justify**

N/A

**Date of registration in national regulatory agency**

24/11/2022

**Primary sponsor**

American University of Beirut

**Primary sponsor: Country of origin**

Lebanon

**Date of registration in primary registry**

24/01/2023

**Date of registration in national regulatory agency**

24/11/2022

**Public title**

Young adults as community mental health workers in humanitarian settings

**Acronym**

**Scientific title**

Young adults as community mental health workers in humanitarian settings: A pilot test of the mechanisms of effect on their own well-being

**Acronym**

**Brief summary of the study: English**

Young adults are resources and agents of change in their communities when they are employed efficiently. Involving young adults in work that influences their communities has positive outcomes for young adults' wellbeing; and for their communities. Still, young adults are often marginalized, particularly in humanitarian settings. If young adults are to be meaningfully engaged as intervention agents, their work should focus on a community need. In the context of a humanitarian crises, has shed light on the need for human resources supporting mental health distress. For this reason, a team of researchers in the United States and Lebanon in partnership with MAPs a Lebanese local NGO, decided to utilize the youth vibrant capacities in Lebanon; targeting Syrian Refugee youths residing in the Beqaa region in particular. Training 25 Young adults (18-24 years of age) on providing an attenuated form of mental health support through the intervention of Problem management plus (PM+) to 400 adults exposed to mental health distress. The aim is to observe the effectiveness of being elements of support to their communities on their own wellbeing, coping mechanisms, and stress. In comparison to being community mental health workers, the team will employ also a group of 20 young adults, who will be trained of tutoring techniques and topics, to deliver tutoring sessions to 200 students from families of the same community and an additional control group of 40 young adults who will only submit survey data.





## Brief summary of the study: Arabic

الشباب هم موارد وعوامل للتغيير في مجتمعاتهم. إن إشراك الشباب في العمل الذي يؤثر على مجتمعاتهم له نتائج إيجابية على رفاهية الشباب؛ ولمجتمعاتهم. ومع ذلك، غالبًا ما يتم تهيمش الشباب، لا سيما في سياق الأزمات الإنسانية. إذا كان الشباب سيشاركون بشكل هادف كوكلاء تدخل يجب أن يرتكز توجيه عملهم على حاجة المجتمع. في سياق الأزمات الإنسانية، الموارد البشرية التي تدعم ضائقة الصحة النفسية لا تكفي، لاجئًا سوريًا شابًا ليكونوا عاملين في مجال الصحة النفسية المجتمعية لتوفير تدخل دعم نفسي واجتماعي 20 سنة عمل هذه الدراسة على تدريب بالغ في مجتمعاتهم. تهدف الدراسة إلى تقييم فعالية كون الشباب عاملين 400 إلى (PM+) منخفض الكثافة يسمى المعالجة المطورة للمشكلات مجتمعيين على عايتهم النفسية و نسب التوتر والتعامل على الشباب أنفسهم من خلال جمع البيانات و عينات الشعور لتحليل نسب الكورتيزول. سيتم طالب في 200 شابًا سيتم تدريبهم لتقديم الدعم التعليمي إلى 20) 1 تقييم هذه النتائج بين العاملين المجتمعيين الشباب ومجموعتين للمقارنة: ( شابًا سيقدّمون بيانات المسح فقط. ستقيم الدراسة أيضًا جدوى ومقبولية الشباب الذين يقدمون الدعم 40) 2 في نفس المجتمع، و (6-1 الصوف النفسي والاجتماعي للبالغين في مجتمعاتهم.

## Health conditions/problem studied: Specify

Managing stress and improving mental health in vulnerable communities through problem management interventions.

## Interventions: Specify

In this pilot study, we aim to evaluate the feasibility, acceptability and fidelity of Syrian refugee young adults as community mental health workers (YA-CMHW), and the impact of this work on outcomes of wellbeing, coping and stress. In addition, we aim to assess the mechanisms leading to any changes in these outcomes.

## Key inclusion and exclusion criteria: Inclusion criteria

Inclusion Criteria: a) being Syrian refugee, (b) being 18-24 years; (c) having completed high school (44); (d) living in one of catchment areas (ITS) in the Bekaa at the time of screening; (e) having been involved in NGOs or service to their community; (f) expressed motivation to serve their community.

## Key inclusion and exclusion criteria: Gender

Both

## Key inclusion and exclusion criteria: Specify gender

## Key inclusion and exclusion criteria: Age minimum

18

## Key inclusion and exclusion criteria: Age maximum

24

## Key inclusion and exclusion criteria: Exclusion criteria

Exclusion criteria: acute mental distress assessed through clinical interview and scales (PHQ9 and PSYCHLOPS) administered by psychiatrist.

## Type of study

Interventional

## Type of intervention

Behavioral treatment

## Type of intervention: Specify type

N/A

## Trial scope

Other

## Trial scope: Specify scope

## Study design: Allocation

Randomized controlled trial

## Study design: Masking

N/A

## Study design: Control

Active

## Study phase

0 (explanatory trials)

## Study design: Purpose

Prevention

## Study design: Specify purpose

N/A

## Study design: Assignment

Parallel

## Study design: Specify assignment

N/A

## IMP has market authorization

## IMP has market authorization: Specify

## Name of IMP

## Year of authorization

## Month of authorization



**Type of IMP**

**Pharmaceutical class**

N/A

**Therapeutic indication**

N/A

**Therapeutic benefit**

N/A

**Study model**

Ecologic or community studies

**Study model: Specify model**

community

**Study model: Explain model**

Observing the effect of delivering social support (low scale mental health intervention) on the mental health, wellbeing of the young adults delivering the intervention rather

**Time perspective**

Prospective

**Time perspective: Specify perspective**

N/A

**Time perspective: Explain time perspective**

We will be following young adults for one year and a half. They will complete a survey 4 times over that time period, and we will collect hair cortisol 6 times over that time period.

**Target follow-up duration**

1

**Number of groups/cohorts**

3

**Biospecimen retention**

Samples without DNA

**Target follow-up duration: Unit**

year

**Biospecimen description**

Hair cortisol samples will be collected to monitor stress level over the period of the study

**Target sample size**

40

**Actual enrollment target size**

**Date of first enrollment: Type**

Anticipated

**Date of first enrollment: Date**

01/02/2023

**Date of study closure: Type**

Anticipated

**Date of study closure: Date**

30/08/2024

**Recruitment status**

Pending

**Recruitment status: Specify**

**Date of completion**



## IPD sharing statement plan

Yes

## IPD sharing statement description

Oversight responsibility: Drs. Afifi and Nakkash (co-PIs) will provide oversight to the trial and the ultimate responsibility for all data safety issues rest with them.

The overall framework for safety monitoring and what information will be monitored: Study data will be accessible at all times to the co-PIs, the site PI (Dr. Ghandour) and to our mental health team. We will be monitoring mental health indicators for the YA-CMHW and ECG groups. The frequency of monitoring, including any plans for interim analysis and stopping rules (if applicable): For both groups (YA-CMHW and ECG), we will be collecting survey data at 4 points

in times but also experience sampling method (ESM) data every three weeks (2-3 questions). This will allow the identification of acute distress or imminent threat of harm to self or others. The data will be reviewed within 24 hours of receipt by our psychiatrist and one of the co-PIs or site PI (Afifi or Nakkash or Ghandour). In addition, the psychologist will be meeting with the YA-CMHW group weekly in group and individual meetings. S/he will assess acute distress or imminent threat of harm to self or others at these meetings. If s/he identifies any CHMW with such outcomes, s/he will immediately refer to MDM, and inform our project psychiatrist and Afifi or Nakkash or Ghandour.

If more than 1/3 of the YA-CMHW exhibit acute distress or imminent threat of harm to self or others at any time during the implementation of the intervention, we will stop this trial. Any adverse or serious adverse event, or unanticipated problems will be reported within 24 hours to the IRB by Dr. Ghandour and to the chair of the DSMB by Dr. Afifi or Nakkash. The project mental health team (psychiatrist and psychologist) will provide immediate guidance on management of the AEs and SAEs. We have a letter of commitment/support from Médecins Du Monde to provide mental health care free of charge to those referred (see letter

of support). Data Safety Monitoring Board (DSMB): We will establish an independent data monitoring and safety board that meets quarterly or more frequently as needed. The board will consist of 5 faculty or staff with expertise in biostatistics and research ethics and/or humanitarian settings; as well as at least 2 community advocates. Three of the faculty/staff board members will be from Lebanon, and 2 from the US; all community advocates will be from Lebanon, and Syrian if possible. The DSMB will also monitor and evaluate the progress of the trial, recruitment of participants, retention, processes and timeliness of data collection, risk and benefits, and other aspects of the progress of the study that might affect outcomes. The DSMB will also highlight external factors that may be important to the progress of our research trials, including concerns about repercussions based on the socio-political situation in Lebanon, or related to scientific advancements relevant to the project.

Adverse events involving risks to participants will be reported by the PIs (Nakkash or Afifi) to the DSMB and to the University's IRB as per their policy. Finally, the DSMB will guide any decisions related to continuation or discontinuing the trials. All the above will, of course, be done with utmost attention to the confidentiality of all trial data.

Additional data URL

Admin comments

Trial status

Approved



## Secondary Identifying Numbers

No Numbers

## Sources of Monetary or Material Support

Name

NIH

## Secondary Sponsors

No Sponsors

## Contact for Public/Scientific Queries

Contact type	Contact full name	Address	Country	Telephone	Email	Affiliation
Public	Lilian Ghandour	Lebanon	Lebanon	+961 3 644 544	lg01@aub.edu.lb	AUB
Scientific	Rima Afifi	USA	Lebanon	+1 (319) 471-5333	rima- affi@uiowa.edu	IOWA university

## Centers/Hospitals Involved in the Study

No Centers/Hospitals

## Ethics Review

Ethics approval obtained	Approval date	Contact name	Contact email	Contact phone
American University of Beirut Medical Center	13/12/2022	Lina Onsi	le08@aub.edu.lb	1-350000 or 1 374374, ext: 5445



## Countries of Recruitment

**Name**

Lebanon

## Health Conditions or Problems Studied

No Problems Studied

## Interventions

Intervention	Description	Keyword
delivering PM+	Effect of delivering PM+ on the mental health and wellbeing of young adults delivering it to members of their community	PM+
delivering tutoring	effect of doing a task (tutoring school kids) on the mental health and wellbeing of young adults delivering it	social intervention

## Primary Outcomes

Name	Time Points	Measure
wellbeing	4 and 6	survey and hair samples
stress	4 and 6	survey and hair samples
resilience	4 and 6	survey and hair samples

## Key Secondary Outcomes

Name	Time Points	Measure
autonomy	4	survey
Engagement	4	survey
Hope	4	survey
meaning	4	survey
accomplishment	4	survey
positive emotions	4	survey
relationships	4	survey



## Trial Results

**Summary results**

**Study results globally**

**Date of posting of results summaries**

**Date of first journal publication of results**

**Results URL link**

**Baseline characteristics**

**Participant flow**

**Adverse events**

**Outcome measures**

**URL to protocol files**