

Determining the effectiveness of a structured group dynamic relaxation-training programme on reducing anxiety and depression symptoms

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| Submission date 01/04/2019 | Recruitment status No longer recruiting | <input type="checkbox"/> Prospectively registered |
| Registration date 08/04/2019 | Overall study status Completed | <input type="checkbox"/> Protocol |
| Last Edited 20/09/2021 | Condition category Mental and Behavioural Disorders | <input type="checkbox"/> Statistical analysis plan |
| | | <input checked="" type="checkbox"/> Results |
| | | <input type="checkbox"/> Individual participant data |

Plain English summary of protocol

Background and study aims

Studies have persistently shown that anxiety disorders cause morbidity, increased use of health care services, decreased work productivity, work absence and will possibly be one of the leading causes of disability in the twenty-first century. In the treatment of anxiety disorders, psychopharmacological interventions with benzodiazepines and selective serotonin reuptake inhibitors (SSRI) have shown their efficacy. Non-pharmacologic interventions may be proposed as an alternative option, with the aim to reduce perceived anxiety and stress and to increase the sense of well being in the general population and relaxation – meditation techniques represent one of the most important alternatives in anxiety intervention worldwide.

The aim of the present study is to determine the effectiveness of an intensive four-week structured group relaxation-training program (sophrology's dynamic relaxation) on anxiety and depression symptoms in primary care patients with moderate and high anxiety levels.

Who can participate?

Patients with moderate and high anxiety levels aged between 18 and 70 years.

What does the study involve?

Participants attend 12 one-hour sessions over a 4 week period. The intervention group will participate in the dynamic relaxation program called "well-being and sophrology" and the control group will participate in a cognitive program based on physical and mental health recommendations.

What are the possible benefits and risks of participating?

Possible benefits are the improvement of anxiety and depression symptoms. No risks expected as a result of participation.

Where is the study run from?

Medical Health Care Centre "Serraparera", Cerdanyola, Barcelona.

When is the study starting and how long is it expected to run for?
May 2015.

Who is funding the study?
Investigator-initiated and funded.

Who is the main contact?
Koen van Rangelrooij
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Contact information

Type(s)
Scientific

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Study information

Scientific Title
Effectiveness of a structured group relaxation-training programme based on sophrology's dynamic relaxation techniques for primary care patients with moderate and high anxiety levels: a randomised controlled trial

Study objectives
The Dynamic relaxation programme (sophrology) is more effective than a physical and mental health recommendations programme in patients with moderate and high anxiety levels.

Ethics approval required
Old ethics approval format

Ethics approval(s)
Approved 20/10/2015, the University Review Board of the Autonomous University of Barcelona (CEIm – Parc de Salut MAR, Dr. Aiguader, 88, 08003, Barcelona; 93 316 06 77; ceic-psmar@imim.es), ref: 2015/6141/l.

Study design
Interventional and experimental study design, single centre, randomized controlled trial,

Primary study design

Interventional

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Anxiety

Interventions

Both programs covered a total of 12 one-hour sessions over 4 weeks (3 sessions a week). The intervention group followed the structured dynamic relaxation program called "well-being and sophrology" and the control group a cognitive program based on physical and mental health recommendations (PMHR). Simple random sampling was used. Two physicians, two nurses and a psychologist from the Medical Health Care Centre guided both the sophrology intervention and the PMHR (Physical and Mental Health Recommendations) control programme.

Intervention Type

Behavioural

Primary outcome(s)

Anxiety was measured using the Hospital Anxiety Depression Scale (HADS) and the State-Trait Anxiety Inventory (STAI) at the beginning of the first and the last session of the training programme.

Key secondary outcome(s)

N/A

Completion date

19/06/2015

Eligibility**Key inclusion criteria**

1. A score >7 for the Hospital Anxiety Depression Scale - anxiety subscale
2. 18 to 70 years of age

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Lower age limit

18 years

Sex

All

Total final enrolment

70

Key exclusion criteria

1. Initiated or changed pharmacological, behavioural or any other therapy during the programme
2. Presented uncontrolled mental illness
3. Planned to participate in other therapies or similar programmes such as Yoga, Mindfulness, meditation, acupuncture, or other
4. During the programme suffered from important stressful life events
5. Not able to participate for linguistic, cultural or physical problems
6. Could not attend sessions regularly (<80% attendance)

Date of first enrolment

04/05/2015

Date of final enrolment

22/05/2015

Locations**Countries of recruitment**

Spain

Study participating centre**Medical Health Care Centre "Serraparera"**

Carrer Diagonal, s/n,
Cerdanyola del Valles

Spain

08290

Sponsor information**Organisation**

Autonomous university Barcelona

ROR

<https://ror.org/052g8jq94>

Funder(s)

Funder type

Other

Funder Name

Investigator initiated and funded

Results and Publications

Individual participant data (IPD) sharing plan

The data sharing plans for the current study are unknown and will be made available at a later date.

IPD sharing plan summary

Data sharing statement to be made available at a later date

Study outputs

| Output type | Details | Date created | Date added | Peer reviewed? | Patient-facing? |
|---------------------------------|---------|--------------|------------|----------------|-----------------|
| Results article | | 01/09/2020 | 20/09/2021 | Yes | No |