Efficacy of text-messages in helping former patients (previously treated in addiction centers) to maintain their substance use goal

Submission date	Recruitment status Stopped	[X] Prospectively registered		
10/07/2017		☐ Protocol		
Registration date	Overall study status	Statistical analysis plan		
12/07/2017	Stopped	☐ Results		
Last Edited	Condition category	Individual participant data		
04/08/2021	Mental and Behavioural Disorders	Record updated in last year		

Plain English summary of protocol

Background and study aims

Text-messaging-based aftercare has been shown to be effective for various mental health problems such as eating disorders and alcohol withdrawal. Former patients can generally be reached easily with text messages, which might help to reduce their risk of relapsing. The aim of this study is to assess the effectiveness of text messages in helping former patients (previously treated in addiction centers) to maintain their substance use goal during the 6 months following inpatient treatment.

Who can participate?

Patients aged 18 and over who have completed their addiction treatment

What does the study involve?

Participants are randomly allocated to one of two groups shortly before leaving the treatment center. Participants in one group receive automatically generated supportive text-messages to help them to avoid relapses. Participants in the other group receive semi-automatically generated, individualised supportive text messages and also telephone calls from a counselor. Text messages are sent for a period of six months after patients have left treatment, first weekly, then biweekly. Both groups are assessed for their maintenance of abstinence or controlled drinking (depending on personal goals) during the 6 months following inpatient treatment.

What are the possible benefits and risks of participating?

Former patients who receive text message based aftercare might be encouraged to avoid relapses. No risks are expected.

Where is the study run from?

- 1. Swiss Research Institute for Public Health and Addiction ISGF (Switzerland)
- 2. Akzente Prävention und Suchttherapie, Luzern (Switzerland)
- 3. Die Alternative, Ottenbach (Switzerland)
- 4. Freihof Küsnacht, Küsnacht (Switzerland)

- 5. Forelhaus, Zürich (Switzerland)
- 6. RehabilitationsZentrum Lutzenberg, Lutzenberg (Switzerland)
- 7. Suchttherapiebärn, Bern (Switzerland)

When is the study starting and how long is it expected to run for? September 2015 to July 2020

Who is funding the study? Federal Office of Public Health (FOPH) (Switzerland)

Who is main contact? Mrs Susanne Schaaf schaaf@isgf.uzh.ch

Contact information

Type(s)

Scientific

Contact name

Mrs Susanne Schaaf

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Contact details

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Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers

PEG201410

Study information

Scientific Title

Efficacy of text messaging-based aftercare following inpatient addiction treatment - DASA (Drug Alcohol SMS Aftercare): a randomised controlled trial

Acronym

DASA – Drug Alcohol SMS Aftercare

Study objectives

Individualised SMS-based intervention (including information on personal reward suggestions and coping strategies on how to deal with tempting situations, as well as in addition phone calls from therapists) is more effective than non-individualised SMS-based intervention to maintain abstinence or controlled drinking (depending on the personally set goal).

Ethics approval required

Old ethics approval format

Ethics approval(s)

Ethics Committee Zurich (lead committee), 19/06/2017, ref: PB_2016_02180

Study design

Multicentre randomised active-controlled parallel-group trial

Primary study design

Interventional

Secondary study design

Randomised controlled trial

Study setting(s)

Other

Study type(s)

Prevention

Participant information sheet

See additional files

Health condition(s) or problem(s) studied

Relapse prevention for former patients who completed addiction treatment successfully

Interventions

Randomisation will be undertaken centrally by the Swiss Reasearch Institute for Public Health an Addiction ISGF. The questionnaires will be handed out to the participating treatment centers, and will be stratified by gender and main problem substance of patients, resulting in four combinations: a) female, alcohol, b) male, alcohol, c) female, drug (not-prescribed opiates, cocaine or cannabis), c) male, drug (ditto). Within each combination, the questionnaires are - according to the randomisation list - assigned to either control group or intervention group. The therapist fills in the questionnaire together with the client. The allocation to the study group will be disclosed on the last page of the questionnaire

The patients will be randomly assigned to one of two study groups shortly before leaving the treatment center:

- 1. Minimal/standardised intervention control group (CG): automatically generated, standardised text-messages, standardised supportive messages: first weekly, then biweekly SMS regarding maintenance of abstinence or controlled alcohol use respectively; support from advisor on demand
- 2. Intervention group (IG): semi-automatically generated, individualised text-messages: first weekly, then bi-weekly SMS regarding maintenance of abstinence resp. controlled alcohol use; individualised supportive text-messages reward, coping strategies) and active contacting by advisor in case of not-maintaining abstinence resp. controlled alcohol use.

SMS messages will be sent for a period of six months after patients have left treatment, first weekly, then biweekly.

The primary outcome measures are assessed at 6-months follow up. Frequency of use of the respective main problem substance within the previous six months will be assessed. Participants indicating that they never used the substance within the previous six months are considered as abstinent (prolonged abstinence). Concerning controlled drinking, the participants should indicate whether or not they always maintained their personal goal of alcohol use within the previous six months (prolonged maintenance of drinking goal).

Intervention Type

Other

Primary outcome measure

Maintenance of abstinence (illegal substances such as opiates, cocaine, cannabis; legal substance: alcohol) or maintenance of controlled drinking (depending on personal goals) during the 6 months following inpatient treatment, assessed by self-report at 6-months follow up

Secondary outcome measures

- 1. Retention to intervention program (intervention group vs control group), assessed by measuring the proportion of participants which did not actively discontinue programme participation during the study period of 6 months by informing the study center via phone, text message or e-mail
- 2. 7-day and 30-day point prevalence abstinence and maintenance of controlled drinking, respectively, assessed by self-report at 6-months follow up
- 3. Acceptance of intervention program by former patients (intervention group vs control group), assessed using the log files of the text messaging system during the study period of 6 months

Overall study start date

07/09/2015

Completion date

31/07/2020

Reason abandoned (if study stopped)

Objectives no longer viable

Eligibility

Key inclusion criteria

- 1. Age >=18 years
- 2. Residential treatment will presumably be finished within one week
- 3. Main problem substances at beginning of treatment: opioids, cocaine, cannabis or alcohol
- 4. Personally set goals: abstinence (re. illegal substances and alcohol) or controlled drinking respectively
- 5. Owning a mobile phone

Participant type(s)

Other

Age group

Adult

Lower age limit

18 Years

Sex

Both

Target number of participants

554

Key exclusion criteria

Cognitive or linguistic deficits or health impairments which impede following the SMS dialogue

Date of first enrolment

01/08/2017

Date of final enrolment

15/06/2019

Locations

Countries of recruitment

Switzerland

Study participating centre RehabilitationsZentrum Lutzenberg

Lutzenberg Switzerland 9426

Study participating centre Suchttherapiebärn

Bern Switzerland 3006

Study participating centre Akzent Prävention und Suchttherapie

Luzern Switzerland 6003

Study participating centre Die Alternative

Ottenbach Switzerland 8913

Study participating centre Freihof Küsnacht

Küsnacht Switzerland 8700

Study participating centre Forelhaus

Zürich Switzerland 8003

Sponsor information

Organisation

Swiss Research Institute for Public Health and Addiction ISGF

Sponsor details

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Sponsor type

Research organisation

Website

www.isgf.ch

ROR

https://ror.org/02crff812

Funder(s)

Funder type

Government

Funder Name

Federal Office of Public Health (FOPH)

Results and Publications

Publication and dissemination plan

Planned publication in a high-impact peer reviewed journal in 2021.

Intention to publish date

31/07/2021

Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study are not expected to be made available due to a lack of participants' consent to share data with other institutions.

IPD sharing plan summary

Not expected to be made available

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Participant information sheet		12/07/2017	12/07/2017	No	Yes