

Reducing no-shows in outpatient forensic mental health care

Submission date 19/11/2024	Recruitment status No longer recruiting	<input type="checkbox"/> Prospectively registered <input checked="" type="checkbox"/> Protocol
Registration date 04/12/2024	Overall study status Ongoing	<input type="checkbox"/> Statistical analysis plan <input type="checkbox"/> Results
Last Edited 02/12/2024	Condition category Other	<input type="checkbox"/> Individual participant data <input checked="" type="checkbox"/> Record updated in last year

Plain English summary of protocol

Background and study aims

The issue of patient non-attendance or tardy cancellation of appointments is a prevalent challenge for healthcare facilities. These occurrences, commonly labeled as 'no-shows', have significant implications for financial costs, treatment progression, and patient waiting periods. This study aims to build a predictive model for no-show for patients in forensic mental health, to implement the model in clinical practice, and to determine if phone call reminders, for the group with a considerable predicted risk of no-show at the next appointment, lead to a reduction in no-show occurrences.

Who can participate?

All appointments for participants who received outpatient treatment at Transfore.

What does the study involve?

A no-show prediction model for outpatient forensic mental health care will be developed. This model will be implemented and tested in clinical practice using data. The six outpatient forensic locations of Transfore, part of the Dimence Groep, will be divided into two different groups. One group, consisting of two locations (Enschede and Zwolle), will implement the model and patients with a high risk of missing their appointment will receive a reminder phone call. These reminder calls are made two days in advance. In the other group, no information is shared about patients with a higher risk of no-show and no intervention is applied.

What are the possible benefits and risks of participating?

Participants may receive a reminder phone call with the opportunity to reschedule their appointment.

A possible risk is that the patient receives an unjustified reminder phone call.

Where is the study run from?

The Dimence Groep, Institute of Mental Health

When is the study starting?

September 2022 to December 2025. 14/09/2022 - Start date for building the predictive model.
25/11/2024 - Start date for testing the model in clinical practice.

Who is funding the study?

The Dimence Groep

Who is the main contact?

Erik de Groot, e.degroot@dimencegroep.nl

Contact information

Type(s)

Public, Scientific, Principal Investigator

Contact name

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Contact details

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Additional identifiers

EudraCT/CTIS number

Nil known

IRAS number

ClinicalTrials.gov number

Nil known

Secondary identifying numbers

DimenceGroep2024NoShow

Study information

Scientific Title

Predictive modelling of no-show in outpatient forensic mental health care: a machine-learning approach.

Acronym

REDUCING

Study objectives

Reminder phone calls to patients with a high predicted risk of no-show at their next appointment will significantly reduce the number of no-show occurrences in outpatient forensic mental health care.

Ethics approval required

Ethics approval required

Ethics approval(s)

Approved 14/11/2024, Scientific Research Committee of Dimence (Burg. Roelenweg 9, Zwolle, 8021EV, Netherlands; +31 038 456 58 51; onderzoek@dimencegroep.nl), ref: CWO/EdG112024

Study design

Interventional sample study

Primary study design

Interventional

Secondary study design

Non randomised study

Study setting(s)

Home, Medical and other records, Other therapist office, Telephone

Study type(s)

Treatment

Participant information sheet

No participant information sheet available

Health condition(s) or problem(s) studied

Preventing no-shows in patients in outpatient forensic mental health care.

Interventions

Patients whose appointments are recorded as not cancelled at the time of the calls and whose predicted risk of no-show is at least 0.50 receive a reminder phone call. These reminder calls are made two days in advance. Reminder calls scheduled on Saturday and Sunday are combined with those on Friday. A call script is developed to ensure uniformity in reminder phone calls. A successful phone call is defined as one in which the caller converses with the patient regarding the appointment with a maximum of two attempts. The outcomes related to the appointment reminder (i.e., whether the patient is reached, appointment cancellation or rescheduling, appointment attendance) are registered.

Intervention Type

Behavioural

Primary outcome measure

Patient attendance at an appointment, defined as 1) attendance and 2) no-show (the patient not attending the appointment or the patient cancelling the appointment less than 24 hours before the appointment), measured using data collected in the clinic's electronic health record system one day after the appointment

Secondary outcome measures

There are no secondary outcome measures

Overall study start date

14/09/2022

Completion date

31/12/2025

Eligibility

Key inclusion criteria

All appointments for participants who received outpatient treatment at Transfore between the earlier mentioned period were included.

Participant type(s)

Patient

Age group

Adult

Lower age limit

18 Years

Sex

Both

Target number of participants

2858 appointments

Key exclusion criteria

Exclusion criteria were appointments that were cancelled by the therapist or the patient more than 24 hours before the appointment and crisis appointments.

Date of first enrolment

25/11/2024

Date of final enrolment

01/06/2025

Locations

Countries of recruitment

Netherlands

Study participating centre

Dimence Groep

Nico Bolkesteinlaan 1

Deventer
Netherlands
7416 SB

Sponsor information

Organisation

Dimence

Sponsor details

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Sponsor type

Hospital/treatment centre

Website

<https://www.dimencegroep.nl/>

ROR

<https://ror.org/010jxjq13>

Funder(s)

Funder type

Hospital/treatment centre

Funder Name

Dimence Groep

Results and Publications

Publication and dissemination plan

Planned publication in a peer-reviewed journal.

Intention to publish date

31/12/2026

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 the current data policy of the study organisation.

IPD sharing plan summary

Not expected to be made available

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Protocol file	version 1.0	20/02/2024	22/11/2024	No	No