

Programme on Adherence to Medication (PAM) randomised controlled trial

Submission date 28/10/2019	Recruitment status No longer recruiting	☒ Prospectively registered ☒ Protocol
Registration date 09/12/2019	Overall study status Completed	☐ Statistical analysis plan ☐ Results
Last Edited 20/07/2026	Condition category Circulatory System	☐ Individual participant data ☒ Record updated in last year

Plain English summary of protocol

Current plain English summary as of 01/12/2023:

Background and study aims

About 9.5 million people in England and Wales receive treatment for hypertension (high blood pressure) from their general practice. However, many people with hypertension and associated conditions (such as diabetes, heart disease and stroke) do not take their medication as prescribed. Not taking medication reduces how well such treatment works and increases healthcare costs; for example, unused medications cost the NHS in England several hundred million pounds per year.

Primary care practitioners have an important role in supporting patients to adhere to their prescribed medication. However, they have limited time to provide ongoing support for adherence, and their time is expensive.

A potential solution is for practitioners to deliver a very brief intervention during a consultation and to use a digital intervention such as text messaging, smartphone app or web-based intervention to support subsequent adherence. Digital interventions have several advantages: they can be fully automated; provide information that is highly tailored to the individual; be interactive; be available at any time; deliver support in real time; deliver support with high fidelity; and be easily updated. A two-component intervention comprising a very brief intervention from a primary care practitioner plus a digital intervention would be inexpensive to deliver, scalable and potentially cost-effective.

About 90% of UK adults and 67% of those aged 65 years and over use a smartphone, and 96% and 87% respectively use a mobile phone capable of text messaging.

We have developed an intervention comprising a very brief intervention delivered remotely by a practice nurse or healthcare assistant in primary care followed by a digital intervention (text messaging programme or smartphone app): the PAM (Programme on Adherence to Medication) intervention. We have shown the intervention to be acceptable, feasible and potentially effective in supporting medication adherence and reductions in blood pressure in patients prescribed treatment for hypertension, as an adjunct to usual care consultations.

The PAM trial will test whether the intervention helps people take their medicines as prescribed and reduces their blood pressure (i.e. whether it is effective) compared with usual care, and also whether it provides value for money. If the findings are positive, the intervention could be rapidly implemented in the NHS, with immediate and long-term benefits to patients and to the health service.

Who can participate?

Patients who have a diagnosis of hypertension (high blood pressure), have been prescribed at least one antihypertensive medication, and have poorly controlled blood pressure or gaps in collecting repeat prescriptions.

What does the study involve?

Participants are randomly allocated to the intervention group or the control group. The intervention group receive a very brief intervention (VBI) delivered remotely (by phone or videolink) by a healthcare practitioner, followed by a digital intervention (text messaging or smartphone app) for up to 14 months. The control group receive usual care only. Follow up is at 6 and 12 months. Participants will be asked to measure their own blood pressure, provide urine and blood samples, and complete questionnaires.

What are the possible benefits and risks of participating?

The potential benefits of participating are that participants allocated to the intervention group may find the intervention helpful in supporting them to take their medications as prescribed, which will be beneficial to their health. Participants allocated to the control group will not benefit from the intervention, but they may find that participation in general and measurement in particular (e.g. self-monitored home blood pressure measurement) may help them to take their medication as prescribed and control their blood pressure. Both groups of participants will also be contributing to evidence about the effectiveness and cost-effectiveness of the intervention. We believe that the risks of participating are low. At worst the intervention may not be effective, and some participants may find the measurement procedures burdensome. Participants are free to withdraw from the trial at any time, and procedures are in place to address any issues that arise.

Where is the study run from?

University of Cambridge (UK)

NHS Cambridgeshire and Peterborough Integrated Care Board (UK)

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

October 2020 to September 2024

Who is funding the study?

NIHR Programme Grants for Applied Health Research (UK)

Who is the main contact?

Prof Stephen Sutton, srs34cam@gmail.com

Previous plain English summary:

Background and study aims

Almost 14.5 million people in the UK receive treatment for high blood pressure (hypertension) from their general practice, including many patients with other conditions such as diabetes,

heart disease and stroke. Treatment for high blood pressure usually consists of one or more blood-pressure-lowering medicines, known as antihypertensives. Many patients will also be prescribed other medicines including statins (to lower cholesterol) and glucose-lowering medicines. Taking medication as prescribed can significantly reduce risks, complications and early deaths associated with these conditions. However, many people with these conditions do not take their medication as prescribed. This reduces how well such treatment works, leading to increased heart attacks and strokes. It also means that a lot of medicines are wasted, which costs the NHS several hundred million pounds a year. GPs and nurses can support patients in taking their medication, but they have limited time and their time is expensive. There is, therefore, a need for low-cost interventions that support patients to take their tablets as prescribed. A promising approach is to use digital interventions such as text messaging and smartphone 'apps' that are accessible from mobile phones. Such interventions have several advantages: they can be fully automated; they can provide information that is highly tailored to the individual; they can be interactive; they can be available at any time; they can deliver support in real-time; the content can be easily updated; and, once they have been developed, they are relatively inexpensive to deliver.

The researchers are proposing that a digital intervention could be used in combination with a brief intervention from the practice nurse or healthcare assistant. Practitioners would give brief medication adherence advice and support to patients during consultations such as blood pressure checks and annual reviews. As part of this, they will 'signpost' patients to a digital intervention which would support them to take their medicines between visits to the practice. With the help of patients, practitioners and other experts, and informed by a review of existing evidence, the researchers will develop a new intervention that has two parts: (i) a very brief medication adherence intervention (lasting under five minutes) delivered by a nurse/healthcare assistant, who then 'signposts' the patient to (ii) a digital intervention designed to provide ongoing behavioural support between practice visits. The researchers will test whether the intervention helps people take their medicines as prescribed and reduces their blood pressure (i.e. whether it is effective) compared with usual care, and also whether it provides value for money and utility of health. If the findings are positive, the intervention could be rapidly implemented in the NHS, with immediate and long-term benefits to patients and to the health service.

Who can participate?

Patients who have a diagnosis of high blood pressure (HBP), or comorbidities of HBP, such as type 2 diabetes or high cholesterol, are prescribed at least one antihypertensive medication, and have poorly controlled HBP or gaps in collecting repeat prescriptions

What does the study involve?

Participants are randomly allocated to the intervention group or the control group. The intervention group receive a 1-minute very brief intervention (VBI), followed by a 12-month text message and/or smartphone app intervention. The control group receive usual care only. Follow up is at 12 months.

What are the possible benefits and risks of participating?

The VBI delivered by the health care professionals has been designed to support medication adherence and to register patients into the text messaging or app intervention. The researchers will use this information to make recommendations to health care providers in primary care about how best to support people take their prescribed medications by delivering very brief advice. The text messaging service and app intervention has been designed to reflect research which suggests digital interventions may enhance medication adherence by providing ongoing support following practitioners' consultations. This intervention will be available to provide advice on taking medications and support patients to self-monitor their medication taking. It is

anticipated that this element of the digital intervention will increase patients' satisfaction with the continuous care they receive from the practice. If feasible, this low-cost intervention could reach people even in the most deprived areas. Practitioners may not directly benefit from the study, but if successful, this intervention may benefit people with long-term health conditions and help other practitioners to achieve practice QOF targets. Furthermore, medication non-adherence reduces the effectiveness of treatment and increases the cost to the NHS from hospital admissions, additional consultations, referrals, investigations and medicine wastage. If this scalable intervention is effective it will most probably be a cost-effective intervention for the NHS. This is a behavioural intervention and does not involve the test of any medical device /equipment or drugs. Thus, in the researchers' view, this is a low-risk study.

Where is the study run from?
University of Cambridge (UK)

When is the study starting and how long is it expected to run for?
October 2020 to October 2023

Who is funding the study?
Programme Grants for Applied Research (UK)

Who is the main contact?
Dr Katerina Kassavou, kk532@medschl.cam.ac.uk

Contact information

Type(s)

Public, Scientific, Principal investigator

Contact name

Prof Stephen Sutton

ORCID ID

<https://orcid.org/0000-0003-1610-0404>

Contact details

University of Cambridge
Primary Care Unit
Department of Public Health and Primary Care
East Forvie Building
Cambridge
United Kingdom
CB2 0SR
+44 (0)1223 330594
srs34cam@gmail.com

Additional identifiers

Integrated Research Application System (IRAS)
268471

Protocol serial number

268471 v0.2 20-10-19

Study information

Scientific Title

Randomised controlled trial of a very brief practitioner-delivered intervention plus a digital intervention to support medication adherence in people prescribed treatment for hypertension in primary care: the Programme on Adherence to Medication (PAM) trial

Acronym

PAM

Study objectives

Current study hypothesis as of 01/12/2023:

The aim of this trial is to estimate the effectiveness and cost-effectiveness of the PAM intervention to improve medication adherence and reduce blood pressure compared with usual care, to inform a decision on whether to implement the intervention in primary care

Previous study hypothesis as of 22/10/2020:

This trial aims to assess the effectiveness and cost-effectiveness of the PAM intervention to support blood pressure control and medication adherence in primary care.

Previous study hypothesis:

Is PAM (cost) effective to support adherence to antihypertensive medications and improve blood pressure as an adjunct to primary care?

Ethics approval required

Ethics approval required

Ethics approval(s)

Approved 05/10/2020, Cambridge East Independent Research Ethics Committee (Health Research Authority, 2 Redman Place, Stratford, London, E20 1JQ, United Kingdom; +44 2071048181; CambridgeEast.REC@hra.nhs.uk), ref: 19/EE/0354

Primary study design

Interventional

Study design

Parallel group multi-centre individually randomized controlled trial

Study type(s)

Efficacy

Health condition(s) or problem(s) studied

Supporting medication adherence in patients prescribed treatment for hypertension

Interventions

Current interventions as of 01/12/2023:

Intervention group: A very brief intervention (VBI) delivered remotely (by phone or video call) by a practice nurse or healthcare assistant followed by a digital intervention (text messaging or smartphone app), in addition to usual care.

Control group: Usual care only.

Previous interventions as of 22/10/2020:

Intervention group: usual care PLUS a very brief intervention (VBI) facilitated by a practice nurse or health care assistant followed by a 12-month text messaging programme or smartphone app.
Comparator group: usual care only.

Previous interventions:

The intervention group will receive a 1-minute very brief intervention, followed by a 3 months text message and/or a smartphone app intervention. The control group will receive usual care only.

Follow up will be at 12 months. Randomisation will be stratified by primary care professionals only.

Intervention Type

Behavioural

Primary outcome(s)

Current primary outcome measure as of 01/12/2023:

Systolic blood pressure obtained from self-monitored home blood pressure measurements at 12-month follow up

Previous primary outcome measure as of 22/10/2020:

Blood pressure measured using A&D Upper Arm Blood Pressure Monitor at 12 months follow up

Previous primary outcome measure:

Systolic blood pressure measured using electronic monitoring devices at baseline and at 12 months follow up

Key secondary outcome(s)

Current secondary outcome measures as of 01/12/2023:

1. Medication adherence measured by chemical adherence testing of urine samples, self-report questionnaires and prescription data from practice records, at 12-month follow up
 2. Diastolic blood pressure obtained from self-monitored home blood pressure measurements at 12-month follow up
 3. Full lipid profile and HbA1c for patients with high cholesterol and type 2 diabetes, respectively, at 12-month follow up
 4. Health related quality of life using EQ-5D-5L and resource use questionnaire at 6-month and 12-month follow up
 5. Process evaluation using audio recordings of the baseline consultation, log files recording use of the digital interventions, qualitative interviews with practitioners and patients, and self-report questionnaires at 12-month follow up
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Previous secondary outcome measures as of 22/10/2020:

1. Medication adherence measured by biochemical testing of the urine and by self-reports (i.e. two items from the MARS questionnaire and two additional single-item measures) at 12 months follow up
 2. Full lipid profile and glucose levels for a subsample of patients, measured using blood samples at 12 months follow up
 3. Quality of life measured using EQ-5D-5L, and resource use using a self-reported questionnaire at 12 months follow up
 4. Process evaluation using an audio recording of the VBI, digital log files during the 12-month intervention, interviews and self-reported questionnaires at 12-month follow up
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Previous secondary outcome measures:

1. Medication adherence measured by urine samples, two self-reported items, and MARS at baseline and 12 months follow up
2. Quality of life measured using EQ5-D at baseline and at 12 months
3. Process evaluation using digital log files and audio-files during the 3-months intervention, and self-reported questionnaires at baseline and 12 months

Completion date

30/09/2024

Eligibility

Key inclusion criteria

Current inclusion criteria as of 01/12/2023:

Patients will be included if they satisfy all the following criteria:

1. Has a diagnosis of hypertension (high blood pressure)
2. Have been prescribed at least one antihypertensive (blood pressure lowering) medication
3. Any Blood Pressure reading higher than 140/90 mmHg OR gaps in collecting repeat prescriptions
4. Can understand English and is able to provide informed consent
5. Has a mobile phone and is familiar with sending and receiving text messages

6. The practice nurse or healthcare assistant is not aware of any other reason why the patient should be excluded

Previous inclusion criteria as of 22/10/2020:

Patients will be included if they satisfy all six criteria:

1. Have a diagnosis of hypertension (high blood pressure)
 2. Have been prescribed at least one antihypertensive (blood pressure lowering) medication
 3. Have a most recent blood pressure reading higher than 140/90 mmHg or gaps in collecting repeat prescriptions
 4. Can understand English and is able to provide informed consent
 5. Have a mobile phone and is familiar with sending and receiving text messages
 6. The practice nurse or health care assistant is not aware of any other reason why the patient should be excluded.
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Previous inclusion criteria:

Patients will be included if they:

1. Have a diagnosis of high blood pressure (HBP), or comorbidities of HBP type 2 diabetes cholesterol
2. Are prescribed at least one antihypertensive medication
3. Have poorly controlled HBP or gaps in collecting repeat prescriptions
4. Are able and use mobile phones
5. Have the capacity to provide informed consent

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Sex

All

Total final enrolment

578

Key exclusion criteria

Current exclusion criteria as of 01/12/2023:

Patients will be excluded if:

1. They have BP > 200/100mm Hg or postural hypotension (>20mm Hg systolic drop)
2. They have a diagnosis of dementia or other cognitive difficulties that could affect study participation

3. They have had a recent severe life-threatening event or are under treatment for another long-term health condition (e.g. cancer)
4. They are taking part in another medication adherence intervention or digital intervention for behaviour change
5. Their BP is not managed by their GP practice

Previous exclusion criteria:

Patients will be excluded if they:

1. Have a diagnosis of dementia or other cognitive difficulties that could affect study participation
2. Have had a recent severe life-threatening event or are under treatment for another long-term health condition (e.g. cancer)
3. Take part in another medication adherence and/or digital intervention

Date of first enrolment

01/11/2020

Date of final enrolment

30/03/2022

Locations

Countries of recruitment

United Kingdom

England

Wales

Study participating centre

University of Cambridge

Primary Care Unit

Department of Public Health and Primary Care

East Forvie Building

Cambridge

England

CB2 0SR

Sponsor information

Organisation

University of Cambridge

ROR

<https://ror.org/013meh722>

Organisation

NHS Cambridgeshire and Peterborough Integrated Care Board

Funder(s)

Funder type

Government

Funder Name

Programme Grants for Applied Research

Alternative Name(s)

NIHR Programme Grants for Applied Research, PGfAR

Funding Body Type

Government organisation

Funding Body Subtype

National government

Location

United Kingdom

Results and Publications

Individual participant data (IPD) sharing plan

Current Individual participant data (IPD) sharing plan as of 01/12/2023:

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

Previous IPD sharing statement as of 08/12/2022:

The datasets generated during and/or analysed during the current study are/will be available upon request from the lead and first author Dr Aikaterini Kassavou (katerina.kassavou@gmail.com).

Type of data: primary data for systolic blood pressure and biochemical medication adherence

When the data will become available: after publication in peer-review journals

What access criteria data will be shared including with whom: data will be shared with researchers conducting secondary analysis of RCTs (e.g. meta-analysis).

Previous IPD sharing statement as of 29/11/2021:

The datasets generated during and/or analysed during the current study are/will be available

upon request from the lead and first author Dr Aikaterini Kassavou (kk532@medschl.cam.ac.uk).
Type of data: primary data for systolic blood pressure and biochemical medication adherence
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IPD sharing plan summary

Other

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Protocol article	Participant information sheet	25/02/2026	21/04/2026	Yes	No
HRA research summary			26/07/2023	No	No
Participant information sheet		11/11/2025	11/11/2025	No	Yes
Protocol (preprint)	Study website	25/02/2026	20/03/2026	No	No
Study website		11/11/2025	11/11/2025	No	Yes