BMJ Open

Bedtime versus morning use of antihypertensives for cardiovascular risk reduction (BedMed): protocol for a prospective, randomised, open-label, blinded end-point pragmatic trial

Scott R Garrison,1 Michael R Kolber,1 G Michael Allan,1 Jeffrey Bakal,2 Lee Green,1,3 Alexander Singer,3 Darryl R Trueman,4 Finlay A McAlister,5 Raj S Padwal,5 Michael D Hill,6 Braden Manns,7 Kimberlyn McGrail,8 Braden O’Neill,9 Michelle Greiver,9 Liesbeth S Froentjes,1 Donna P Manca,1 Dee Mangin,10 Sabrina T Wong,10 Cathy MacLean,11 Jessica EM Kirkwood,1 Rita McCracken,12,13 James P McCormack,14 Colleen Norris,15 Tina Korownyk1

ABSTRACT

Introduction Sleep-time blood pressure correlates more strongly with adverse cardiovascular events than does daytime blood pressure. The BedMed trial evaluates whether bedtime antihypertensive administration, as compared with conventional morning use, reduces major adverse cardiovascular events.

Methods and analysis Design

Prospective randomised, open-label, blinded end-point trial.

Participants Hypertensive primary care patients using blood pressure lowering medication and free from glaucoma.

Setting Community primary care providers in 5 Canadian provinces (British Columbia, Alberta, Saskatchewan, Manitoba and Ontario) are mailing invitations to their eligible patients. Social media campaigns (Google, Facebook) are additionally running in the same provinces.

Intervention

Consenting participants are allocated via central randomisation to bedtime vs morning use of all antihypertensives.

Follow-up

(1) Telephone or email questionnaire at 1 week, 6 weeks, 6 months and every 6 months thereafter, and (2) accessing linked governmental healthcare databases tracking hospital and community medical services.

Primary outcome

Composite of all-cause death, or hospitalisation for myocardial infarction/acute-coronary syndrome, stroke or congestive heart failure.

Secondary outcomes

Each primary outcome element on its own, all-cause hospitalisation or emergency department visit, long-term care admission, non-vertebral fracture, new glaucoma diagnosis, 18-month cognitive decline from baseline (via Short Blessed Test).

Select other outcomes

- Recruiting through primary care providers, having minimal exclusion criteria and reducing barriers to participation by communicating directly with participants, helps to ensure accurate data collection and good generalisability to primary care populations.
- Beyond an assessment of efficacy, multiple potential harms are being evaluated.
- Members of the public with hypertension are making substantial contributions to study design and conduct through our 10-member patient working group.
- If we observe relative risk reductions for the primary outcome that are smaller than 17%, it is unlikely we will be able to declare those differences statistically significant with the planned sample size.

Self-reported nocturia burden at 6 weeks and 6 months (no, minor or major burden), 1-year self-reported overall health score (EQ-5D-5L), self-reported falls, total cost of care (acute and community over study duration) and mean sleep-time systolic blood pressure after 6 months (via 24-hour monitor in a subset of 302 sequential participants).

Primary outcome analysis

Cox proportional hazards survival analysis.

Sample size

The trial will continue until a projected 254 primary outcome events have occurred.

Current status

Enrolment ongoing (3227 randomised to date).

Ethics and dissemination

BedMed has ethics approval from six research ethics review boards and will publish results in a peer-reviewed journal.

Trial registration number NCT02990663.
INTRODUCTION

Blood pressure (BP) normally exhibits a circadian rhythm with relatively lower pressures during sleep. Lack of this sleep time ‘dip’ correlates strongly with adverse cardiovascular events such as myocardial infarction (MI), stroke and congestive heart failure (CHF), and BP correlates most strongly with such events when measured at night (ie, during sleep).\(^2\)-\(^5\) Given some antihypertensive medications might lower sleep time BP more effectively when administered at bedtime,\(^6\) administration time could conceivably alter the degree of cardiovascular risk reduction these medications provide.

In 2010, Spanish researchers published the first hypertension trial to compare bedtime with morning antihypertensive administration and examine mortality and morbidity outcomes.\(^7\) The results of this randomised controlled trial (RCT), the MAPEC trial (Monitorizacion Ambulatoria para Prediccion de Eventos Cardiovasculares, i.e. Ambulatory Blood Pressure Monitoring for Prediction of Cardiovascular Events), were striking, reporting a 61% relative reduction in a composite of major adverse cardiovascular events (MACE). Despite the obvious clinical importance of this finding, however, hypertension guidelines have yet to endorse bedtime prescribing.\(^8\)-\(^11\) This presumably relates to concern over irregularities in the reporting of MAPEC’s results and methods.\(^12\)-\(^13\) The MAPEC trial registry, for instance, was attributed to at least eight other RCTs,\(^14\)-\(^21\) making it appear to describe a general programme of research, and not the methods of a single RCT. Following this, in 2019, the same principal investigator published another RCT favouring bedtime over morning antihypertensives, the Hygia trial, which reported a 45% relative reduction in MACE.\(^22\) Again, however, irregularities in the reporting of Hygia’s results and methods, including a lack of clarity over how randomisation and allocation were carried out, has led to calls for independent confirmation of these findings before bedtime prescribing of antihypertensives is embraced.\(^13\)-\(^23\)-\(^25\)

BedMed is a large community-based RCT intended to replicate an MAPEC-like timing intervention in a hypertensive Canadian primary care population. BedMed randomises participants to take all existing BP medication (as tolerated) at bedtime, compared with conventional morning use, and tracks mortality and morbidity using regularly collected administrative health claims and participant self-report. This protocol is prepared in accordance with SPIRIT (Standard Protocol Items: Recommendations for Interventional Trials) guidelines.\(^26\)

Objectives

Main
To determine whether a bedtime versus morning antihypertensive administration time influences mortality or cardiovascular morbidity.

Secondary
To determine whether a bedtime versus morning antihypertensive administration time adversely influences cognitive ability, visual acuity, risk of falls and fractures, or nocturia.

METHODS

Trial design

BedMed is a phase 4 pragmatic clinical trial with an adaptive, event-driven, parallel enrolment, prospective randomised open blinded-endpoint design.\(^27\) Here, ‘adaptive’ refers to the potential future exclusion of new participants whose only antihypertensive is a diuretic, if adherence to bedtime allocation in such individuals is poor (see Adherence to bedtime diuretics substudy).

Recruitment began in March 2017, and the trial will continue until 254 primary outcome events have been observed (the number of events in MAPEC). Based on current ongoing enrollment (3227), and an observed 2.0% annual event rate, final analysis is anticipated in fall 2023.

Setting and recruitment

Pragmatic trials collaborative

Most recruitment (~78%) is through community family physicians (>400) who own and operate independent clinics. These providers are spread widely across five participating provinces (Alberta, British Columbia, Manitoba, Saskatchewan and Ontario), but affiliated with the Pragmatic Trials Collaborative (www.PragmaticTrials.ca), a practice-based research network which is coordinating the trial. Nurse practitioners with their own practice panel (seven at present) are also participating.

Each clinic uses their own electronic medical record to create a list of hypertensive patients and the primary care provider (PCP) removes those they consider palliative or incapable of informed consent. The study team then provides the clinic with recruitment envelopes, which the clinic addresses and mails to these potentially eligible patients. The envelopes contain (1) a letter of introduction from the patient’s PCP and (2) a pamphlet describing the trial and providing contact information (online supplemental files 1 and 2). Interested patients call the study team where research assistants answer questions, determine eligibility and obtain consent either in real-time via email (>80% of participants opt for this) or by letter-mail for handwritten consents.

Social Media

All hypertensive residents of our five participating provinces are eligible for BedMed, whether or not their PCP is involved. While this can happen through word-of-mouth, a social media campaign (Google and Facebook Ads) is being employed to inform the public about the trial. These Ads (online supplemental video 1) direct individuals to a landing page (https://bedmedstudy.ca/)
providing trial information, a check of eligibility and telephone/email contact information for the study team.

**Trial population**

**Inclusion criteria**
- Clinician diagnosis of hypertension (by any physician or nurse practitioner).
- Taking ≥1 BP-lowering medication once daily, or PCP willing to convert ≥1 BP-lowering medication to once daily.
- ≥18 years of age.
- Community-dwelling (i.e., not residing in a nursing home).

**Exclusion criteria**
- Considered palliative or unable to consent by PCP.
- Sleep disrupting work (more than three shifts/month during participant’s regular sleeping hours).
- Glaucoma diagnosis, or using glaucoma medication (safety exclusion: nocturnal hypotension, which bedtime BP meds could worsen, has been associated with optic neuropathy in glaucoma patients).28–30

**Randomisation and allocation**

Consenting participants receive their random allocation to bedtime vs morning BP medications while dialoguing directly with a research assistant who has no preceding clinical interactions with that participant and who obtains their allocation (stratifying by province with random blocks of 10 or 12) from the central REDCap server’s randomisation module, ensuring irreversible and concealed allocation.

**Intervention**

**Treatment**
Use of all once-daily BP-lowering medication(s) at bedtime.

**Control**
Use of all once-daily BP-lowering medication(s) in the morning.

**Implementation**

Participants choose between having their PCP assist their timing change (using the PCP’s judgement on how and what to change), or being assisted by the research assistant with whom they are dialoging. Only PCPs assist with timing changes if participants describe heart disease, or if their BP medications include Tiazac XC or Diltiazem XC (which have delayed-release kinetics), furosemide, isosorbide mononitrate/dinitrate, or alpha blockers (medications whose timing decision may be more complicated). PCPs can convert twice daily medications to once daily alternatives, but this is not actively promoted.

Research assistants only change the timing of once daily medications, with a limit of one medication change per week (using the order ACE inhibitors angiotensin receptor blockers, calcium channel blockers, beta-blockers, diuretic-containing medications, other). They advise participants to make the switch by delaying the next dose until the allocated time, and continuing that schedule. If bedtime use is problematic, they ask participants to try taking their BP meds with dinner. If morning use is problematic, they ask participants to try taking it with lunch. Participants with regularly reversed sleep schedules (i.e., sleeping during the day) take their BP medications when they get up, or when they go to bed, not according to the time of day.

At each follow-up, participants are asked about medication timing, and encouraged to adhere to allocation. No devices to separately monitor adherence are in use. As a memory aid, all participants are advised to place pill bottles near objects they use when transitioning to or from bed (e.g., toothbrush, denture case, alarm clock), or to use an AM/PM dosette. If participants report a new diagnosis of glaucoma, they are advised to take their BP medications in the morning, regardless of allocation, to minimise the risk of optic neuropathy.

**Follow-up and data management**

**Research assistant interactions**

All participant interactions with research assistants are unblinded and recorded directly into the University of Alberta’s implementation of the REDCap data management platform.31 The following interactions are scheduled relative to the date of randomisation.

**Baseline:** Telephone interaction to (1) obtain baseline characteristics, (2) conduct the Short-Blessed Test to assess cognitive function, and (3) randomise the participant. May be split over multiple interactions (participant’s choice).

**One week:** Telephone interaction to troubleshoot timing change problems and encourage adherence.

**Six weeks:** Telephone interaction to gather information on adverse effects and outcomes.

**Six months:** Telephone interaction, or REDCap email survey (participant’s choice), to gather adverse effects and outcomes.

**Twelve months:** Same as 6 months+EQ-5D-5L quality-of-life survey (EuroQol Group’s health-related quality-of-life instrument).

**Eighteen months:** Same as 6 months+follow up Short Blessed Test (but available by telephone only).

**Every 6 months thereafter:** Same as 6 months.

**Administrative claims data**

All Canadian provinces have publicly funded healthcare systems and maintain linkable healthcare databases tracking medical services rendered during healthcare interactions for all their residents. This includes community physician services and diagnoses (whether by specialists or generalists), prescriptions dispensed, reasons for hospitalisation and vital statistics (i.e., mortality). BedMed participants consent to these datasets being accessed and analysed to support the trial, providing both outcomes and baseline characteristics.
Twenty-four-hour ambulatory BP monitoring
To assess between-group differences in achieved BP, we intended to carry out 24-hour BP monitoring on a consecutive sample of 151 intervention and 151 control subjects residing in 6 Alberta communities at 6 months (providing 90% power to detect the difference in overnight systolic BP observed in MAPEC). Although we will be able to reach our intended sample size, the timing of these measurements has been substantially delayed for many participants due to both logistic hurdles, and the COVID-19 pandemic. Participants are provided a copy of their test results, which are also faxed (if they consent) to their PCP.

Outcomes
Unless otherwise stated, all outcomes are recorded over the duration of the study.

Primary
Major adverse cardiovascular events
- Defined as first occurrence of either all-cause death or hospital admission/emergency department (ED) visit for acute coronary syndrome/MI, stroke CHF.

Secondary
1. Each component of the primary outcome individually.
2. All-cause hospitalisation/ED visit.
3. Long-term care (LTC) admission (ie, to nursing home or assisted living facility).
4. Non-vertebral fracture.
5. New glaucoma diagnosis.
6. Cognitive decline at 18 months
   - Defined as ≥ 2-point worsening in cognitive performance compared to baseline, as measured by the Short Blessed Test.

Supplementary safety outcomes
1. Vision
   - Vision self-reported as ‘much worse’ compared with the last follow-up at any point, or ‘slightly worse’ than the last follow-up, on two or more occasions (Note: vision is reported, every 6 months, as either ‘unchanged’, ‘slightly worse’ or ‘much worse’ than the last follow-up).
2. Cognition
   - New ‘impairment consistent with dementia’ at 18 months (Short Blessed Test newly ≥10) or new diagnosis of dementia at any point during follow-up.
3. Symptomatic Hypotension
   - Self-reported light-headedness, or feeling faint without loss of consciousness, in the prior month.
   - Self-reported fainting (loss of consciousness) in the prior month.
   - Self-reported falling in the prior month.
   - Hip fracture.
   (Note: at 6 weeks, 6 months and every 6 months thereafter, participants are separately asked whether they have felt lightheaded, fainted, or fallen in the last month).
4. Nocturia
- Self-reported change from baseline in the number of overnight urinations per week (at 6 weeks and 6 months).
- Self-reported nocturia burden in the prior month, recorded as no nocturia, or nocturia that is ‘no problem’, ‘minor problem’ or ‘major problem’ (at 6 weeks and 6 months).

Cost
1. Acute care costs (estimated from each hospital admission’s resource intensity weight and length of stay)*.
2. Total cost of care (acute care costs+medication costs+physician billings)*.
*All cost measures are derived entirely from administrative claims data, and not from self-report. If claims data is not available for some participants, they will be excluded from this analysis.

Exploratory
1. Self-reported overall health score (via EQ-5D-5L) at 12 months.

Process
1. Proportion of BP medication doses taken at the allocated time at 6 months (two times per day medications being considered as half dose in the AM and half dose in the PM for this calculation)†.
2. Sleep-time systolic BP after 6 months (consecutive sample of 302 Alberta residents)†.
†Although blinded to individual participant process outcomes, investigators are unblinded to the aggregated results for adherence to allocation time, and to the isolated results from the 24-hour BP assessments. This allows for consideration of protocol alterations should the intervention appear poorly applied. Investigators are otherwise fully blinded to all trial outcomes.

End-point adjudication
Administrative data
Administrative data derived outcomes will be identified using established and validated coding algorithms. Physicians providing these diagnoses are generally acute care providers (emergency physicians, hospitalists, specialists) who are unaffiliated with the BedMed trial.

Adjudication panel
Most primary and secondary outcomes are being collected in duplicate (ie, by administrative claims and participant self-reporting of the same events). This information will be reviewed by a panel of three physicians blinded to allocation. If the panel deems both data sources to be concordant, those events will be considered valid, and the event date in administrative claims will be used. When events are discordant (eg, only present in one of the two data sources or differing in diagnoses) the participant’s PCP will be contacted to provide the adjudication panel with more information, including their opinion on whether the event occurred. The exception is all-cause hospitalisation/ED visits, where we will preferentially use only

Garrison SR, et al. BMJ Open 2022;12:e059711. doi:10.1136/bmjopen-2021-059711
administrative claims data, believing it to be highly accurate, and being more challenging to confirm with PCPs given the high number of such occurrences.

**Sample size determination**

BedMed is event driven, and originally sought to observe 406 primary outcome events before stopping. We chose this event target believing this was the largest number of events a network our size could detect with 3 years of observation. However, because patients receiving recruitment packages are less likely to enrol than expected (projected enrolment 12%, actual enrolment 6%), and because the overall annual event rate is at the low end of expectations (2.0%), we have reduced our event target for stopping to 254, which matches the number of events observed in MAPEC. Assuming meaningful covariates, 254 events should allow observed risk ratio differences of ~17% or larger to be declared statistically significant. To estimate when this number of events has likely been reached, Alberta Health Services is tracking the primary outcome event rate in Alberta BedMed participants on a quarterly basis. We then extrapolate this to the trial as a whole using the number of participant years of observation. At the current rate of events and enrolment, BedMed should conduct its final analysis in fall 2023.

**Statistical analysis**

**Intention-to-treat assumptions**

*Lost to follow-up*

If participants are lost to follow-up, but medical services continue to be recorded within administrative claims data, we will treat them as though they were still active in the study and censor survival data on the last date of medical services, or indication of death, whichever occurs later. If no such medical claims exist, data will be censored on the last day of successful telephone or email follow-up.

*Withdrawal*

Participants withdrawing from the study are asked to allow us to continue to follow their administrative claims data. If they agree (as the majority do), we will continue to use administrative claims outcomes for those individuals as per the loss to follow-up description. If they do not agree, survival data will be censored on the date of withdrawal.

*Missing data*

For each analysis, we will either impute a value from subsequent or preceding follow-up visits, or exclude a participant from analysis. How we deal with missing data will be specific to each analysis and prioritise either minimising bias, or being conservative when bias is unavoidable (ie, biasing against benefit and towards harm, for the intervention).

*Non-adherence*

Non-adherence to allocation will not exclude participants from analysis unless the outcome of interest is a harm that only makes sense to assess while on-treatment (eg, assessing how nocturia differs in diuretic users switched to bedtime, compared with non-diuretic users making the same switch).

**Selecting regression covariates**

Analyses of dichotomous outcomes will use a maximum of 1 covariate per 10 outcomes, and analyses of continuous outcomes will use a maximum of 1 covariate per 20 randomised subjects. The covariate list for each analysis is predefined in table 1, and all are measured at baseline. We will always use the maximum number of covariates possible, selected in the order given (ie, we will not undertake stepwise addition or subtraction).

**Subgroup analyses**

We will repeat the primary outcome analysis for those with and without the following baseline characteristics: age ≥75, sex, physically frail (score ≥3 on physical frailty subscale of the Tilburg questionnaire), polypharmacy (≥5 medications), Overall Health Score ≤75, resistant hypertension (≥3 BP-lowering medications), CHF, diabetes, CAD (coronary artery disease), stroke or TIA, sleep apnoea, chronic kidney disease (with or without dialysis), sedentary (exercise < 0 days per week).

**Sensitivity analyses**

We will present, according to treatment group, the baseline characteristics of those whose data was censored due to withdrawal or lost to follow-up, and compare these characteristics to those who were not censored in this way using Fisher’s exact test.

**Patient and public involvement**

**Patient working group**

BedMed has a 10-member patient working group helping to guide the trial. The group began meeting in 2016 prior to any recruitment to review and revise (1) recruitment materials, (2) phrasing of questions and (3) outcomes to be collected through self-report. Working group members have also assisted in hiring research staff, in further revising recruitment materials mid-study to increase enrolment, and in constructing a social media campaign. We anticipate working with our patient partners to make decisions, if needed, following our interim analysis in spring 2022, to interpret final results in 2023, and to help disseminate findings.

**Patient-driven substudy**

The draft BedMed protocol was presented in 2015 to a group of ~25 seniors prior to study registration and grant application. Feedback from this presentation resulted in the substudy to determine whether diuretics can be taken at bedtime without troublesome nocturia threatening adherence.

**SUBSTUDIES**

**Adherence to bedtime diuretics**

Diuretics are widely believed to promote nocturia if taken later in the day, and are typically recommended for...
morning use only as a result.\textsuperscript{35, 36} However, this recommendation is largely opinion based. Whether or not participants will adhere to bedtime diuretic dosing is unclear. To determine this, we will examine, at 6 weeks and 6 months, self-reported nocturia burden (no, minor, major), number of overnight urinations per week, and adherence to bedtime allocation, in the first 203 AM diuretic-only users randomised to bedtime and being followed for 6 months, and compare this to all those switching a single AM non-diuretic to bedtime during the same period.

Table 1: Analysis plan

| Outcome | Method | Covariates |
|---------|--------|------------|
| **Primary** | | |
| Major adverse cardiovascular events | Cox proportional hazards | Age, sex, frailty score\textsuperscript{*}, current smoker, no of non-BP medications, Overall Health Score\textsuperscript{†}, prior 6 months hospitalisation, CHF, diabetes, CAD, stroke or TIA, CKD\textsuperscript{†}, dialysis, BMI >35, BMI <20, sleep apnoea, exercise days\textsuperscript{§}, province (four variables) |
| **Secondary** | | |
| All-cause mortality | Cox proportional hazards | Age, frailty score\textsuperscript{*}, no of non-BP medications, Overall Health Score\textsuperscript{†}, prior 6 months hospitalisation, CHF, diabetes, CAD, CKD\textsuperscript{†} |
| Hospitalisation for stroke | Cox proportional hazards | Age, stroke or TIA, CAD, current smoker, sex, diabetes, exercise days\textsuperscript{§}, BMI >35 |
| Hospitalisation for CHF | Cox proportional hazards | Age, CHF, CAD, diabetes, CKD\textsuperscript{†} |
| All-cause hospitalisation/ED visit | Cox proportional hazards | Age, sex, frailty score\textsuperscript{*}, current smoker, no of non-BP medications, Overall Health Score\textsuperscript{†}, prior 6 months hospitalisation, CHF, diabetes, CAD, stroke, TIA, Short Blessed Test score, CKD\textsuperscript{‡}, dialysis, BMI >35, BMI <20, COPD, province |
| Non-vertebral fracture | Cox proportional hazards | Age, Overall Health Score\textsuperscript{†}, BMI, no of non-BP medications, frailty score\textsuperscript{*}, stroke (not TIA), sex, CHF, exercise days\textsuperscript{§}, TIA, prior 6 months hospitalisation |
| LTC admission | Cox proportional hazards | Age ≥80, Short Blessed Test score, frailty score\textsuperscript{*} |
| New glaucoma diagnosis | Cox proportional hazards | Age, diabetes, CAD or stroke or TIA, CHF, COPD, CKD\textsuperscript{‡}, sleep apnoea, BMI, exercise days\textsuperscript{§}, Short Blessed Test Score |
| 18-month cognitive decline | Poisson regression | Age, sex, frailty score, no of non-BP medications, Overall Health Score\textsuperscript{†}, CHF, stroke, TIA, COPD, BMI, exercise days\textsuperscript{§}, province |
| **Supplementary safety** | | |
| Worsening of vision | Poisson regression | Age, diabetes, CAD or stroke or TIA, CHF, COPD, CKD\textsuperscript{‡}, Overall Health Score\textsuperscript{†} |
| New impairment consistent with dementia | Poisson regression | Age, sex, frailty score, no of non-BP medications, Overall Health Score\textsuperscript{†}, CHF, stroke, TIA, COPD, BMI, exercise days\textsuperscript{§}, province |
| Light-headedness in last Month | Poisson regression | Age, frailty score, no of non-BP medications, Overall Health Score\textsuperscript{†}, CHF, stroke, TIA, sex, exercise days\textsuperscript{§}, BMI, province |
| Syncope in last month | Poisson regression | Age, frailty score, no of non-BP medications, Overall Health Score\textsuperscript{†}, CHF, stroke, TIA, sex, exercise days\textsuperscript{§}, BMI, province |
| Falling in last month | Cox proportional hazards | Age, Overall Health Score\textsuperscript{†}, BMI, no of non-BP meds, frailty score, stroke (not TIA), sex, CHF, exercise days\textsuperscript{§} |
| Hip fracture | Cox proportional hazards | Age, Overall Health Score\textsuperscript{†}, BMI, no of non-BP medications, frailty score, stroke (not TIA), sex, CHF, exercise days\textsuperscript{§} |
| Change in overnight urinations/week | Mann-Whitney or t-test | N/A |
| Nocturia a major burden | Fisher’s exact test | N/A |
| **Cost** | | |
| Acute care costs | Multiple linear regression | Age, sex, frailty score, current smoker, no of non-BP medications, Overall Health Score\textsuperscript{†}, prior 6 months hospitalisation, CHF, diabetes, CAD, stroke, TIA, Short Blessed Test score, CKD\textsuperscript{‡}, dialysis, BMI >35, BMI <20, COPD, province |
| Total cost of care | Multiple linear regression | Age, sex, frailty score, current smoker, no of non-BP medications, prior 6 months hospitalisation, CHF, diabetes, CAD, stroke, TIA, Short Blessed Test score, CKD\textsuperscript{‡}, dialysis, BMI >35, BMI <20, COPD, province |

*Score on physical frailty subscale of the Tilburg questionnaire; continuous 0–8.
†From EQ-5D-5L; continuous 0–100.
‡Not including dialysis.
§‘How many days in the past week have you exercised for 30 min or more, vigorously enough to raise your breathing rate?’; continuous 0–7.
ACS, Acute Coronary Syndrome; BMI, body mass index; BP, blood pressure; CAD, Coronary Artery Disease; CHF, congestive heart failure; CKD, Chronic Kidney Disease; COPD, Chronic Obstructive Pulmonary Disease; ED, emergency department; EQ-5D-5L, EuroQol Group’s health-related quality-of-life instrument; LTC, long-term care; MI, myocardial infarction; N/A, not applicable; TIA, Transient Ischaemic Attack.
Assuming equal numbers in both groups, and 75% adherence to allocation time in non-diuretic users, this should provide 90% power to detect a 20% relative reduction in adherence in diuretic users.

Volunteer bias
Concern has been raised that randomised trial participants are poorly representative of real world populations. We will examine, using Alberta administrative claims, how baseline characteristics and preventive health behaviours differ in four distinct Alberta populations: (1) All BedMed-eligible patients attached to participating PCPs, (2) BedMed participants who enrolled after a PCP-letter, (3) BedMed participants responding to social media advertisement and (4) All BedMed-eligible Albertans. We will compare (1) Demographics (age, sex, postal code derived deprivation index, rural residence), (2) Comorbidities (diabetes, CAD, stroke, osteoarthritis, CHF, chronic obstructive pulmonary disease, dementia, hip fracture, CKD, dialysis, hospital admission in prior 6 months plus accompanying length of stay and resource intensity weighting), (3) Preventive therapies (prior 3 years shingles vaccine, statin use, osteoporosis medication), (4) Screening tests (prior 3 years PAP smear, colonoscopy, mammogram, FIT testing, PSA testing) and (5) clinical outcomes postrandomisation (death, BedMed primary outcome, all-cause hospitalisation or ED visit along with length of stay and resource intensity weighting, nursing home admission, new glaucoma diagnosis/treatment / surgery, hip fracture, and new dementia diagnosis). To substitute for the date of randomisation, we will use the date of PCP mailout for BedMed-eligible PCP-attached patients, and the date providing the same mean number of years of observation for all BedMed-eligible Albertans.

‘Nudge sentence’ recruitment strategy
Two years into recruiting, we hypothesised that altering the physician letter of introduction to state that a large number of people were already participating might improve the response rate. Online supplemental file 3 shows the new physician letter. The added wording states: ‘This study already has over 1700 Canadians with high blood pressure taking part. If you too choose to participate...’. As of March 2019, providers are given an equal number of both recruitment envelopes, sealed and shuffled together, for them to address and mail. Both letters are otherwise identical save for the date on the letter (odd numbered for the new version, even numbered for the original). Participants calling to enrol are asked the date on the letter to determine which version they are responding to, allowing a pseudorandom assessment of the ability of such a ‘nudge’ sentence to improve enrolment. This substudy will continue until recruitment ends, with sample size determined by the number of letters mailed during that interval.

EARLY STOPPING
Independent data safety monitoring board
Outcomes from all provinces will be collected at the end of 2021. Each analysis described in this protocol will then be carried out, and presented to the Cochrane Hypertension Working Group (our independent data safety monitoring board, IDSMB).

Stopping Rules: If p is ≤0.001 for primary outcome benefit (the Haybittle-Peto boundary), or if p is ≤0.05 for harm, the IDSMB will apply clinical judgement and decide whether to recommend to the principal investigator that the trial be stopped early.

Competing studies
A trial similar to ours, the UK’s TIME trial, will likely release results ahead of BedMed. If convincing benefit is demonstrated, we will ask our IDSMB to weigh this new information and consider again whether early stopping is recommended.

Our group is also conducting a separate RCT of the same antihypertensive timing intervention in hypertensive LTC residents (BedMed-Frail). As both trials share the same IDSB, interim data from both trials could be weighed in early stopping discussions for either trial.

DISSEMINATION
Results will be published in a peer-reviewed journal, and summarised in knowledge translation vehicles targeted at PCPs, and the general public. We will also invite trial participants to a results webinar where they can directly pose questions to the principal investigator.

Author affiliations
1Family Medicine, University of Alberta, Edmonton, Alberta, Canada
2Provincial Research Data Services, Alberta Health Services, Edmonton, Alberta, Canada
3Family Medicine, University of Manitoba College of Medicine, Winnipeg, Manitoba, Canada
4BedMed Patient Working Group, Edmonton, Alberta, Canada
5Medicine, University of Alberta, Edmonton, Alberta, Canada
6Clinical Neurosciences, University of Calgary, Calgary, Alberta, Canada
7Nephrology, University of Calgary, Calgary, Alberta, Canada
8Centre for Health Services and Policy Research, University of British Columbia, Vancouver, British Columbia, Canada
9Family and Community Medicine, University of Toronto, Toronto, Ontario, Canada
10Family Medicine, McMaster University, Hamilton, Ontario, Canada
11Academic Family Medicine, University of Saskatchewan, Saskatoon, Saskatchewan, Canada
12Family Medicine, Providence Health Care, Vancouver, British Columbia, Canada
13Family Practice, University of British Columbia Faculty of Medicine, Vancouver, British Columbia, Canada
14Faculty of Pharmaceutical Science, University of British Columbia Faculty of Dentistry, Vancouver, British Columbia, Canada
15Faculty of Nursing, University of Alberta, Edmonton, Alberta, Canada

Twitter Kimberlyn McGrail @kimchspr, Dee Mangin @DeeMangin and Rita McCracken @DrRitaMc

Contributors BedMed was conceived and designed by SRG with input from MRK, GMA, JB, LG, AS, FAM, RSP, MDH, BM, KM, DPM, STW, RM, JPM, CN and TK. Each of these authors also participated as coapplicants on the grants that fund the trial with SRG as the nominated principal applicant. Family physicians were recruited by SRG with help from MRK, GMA, TK, AS, BO’N, MG, DPM, DM, CM, STW
Open access
and JMK. DRT participates on the BedMed Patient Working Group and helped to revise recruitment materials, and to design the social media campaign. SRG and JB created the analysis plan. JB, SRG, AS, FAM and KM are coordinating access to administrative claims data. SRG and LSF wrote the draft manuscript, with all authors providing critical feedback.

**Funding** BedMed is funded by a Support for Patient Oriented Research (SPOR) Innovative Clinical Trial Multi-Year Grant (REF#: 151212) from the Canadian Institutes of Health Research (CIHR), and from a Partnership for Research and Innovation in the Health System (PRIHS) Grant from Alberta Innovates (REF#: 201 500 912 G201900045). It also receives in-kind research assistance from Enhancing Alberta Primary Care Research Networks (ErAC), which itself is funded entirely by Alberta Innovates, and received pilot funding from the Northern Alberta Family Medicine Fund (University of Alberta Dept of Family Medicine funding).

**Disclaimer** Funders have no role in, or authority over, trial design, data collection, management, analysis, interpretation, writing, or decisions to submit for publication.

**Competing interests** None declared.

**Patient consent for publication** Not applicable.

**Provenance and peer review** Not commissioned; peer reviewed for ethical and funding approval prior to submission.

**Supplemental material** This content has been supplied by the author(s). It has not been vetted by BMJ Publishing Group Limited (BMJ) and may not have been peer reviewed. Any opinions or recommendations discussed are solely those of the author(s) and are not endorsed by BMJ. BMJ disclaims all liability and responsibility arising from any reliance placed on the content. Where the content includes any translated material, BMJ does not warrant the accuracy and reliability of the translations (including but not limited to local regulations, clinical guidelines, terminology, drug names and drug dosages), and is not responsible for any error and/or omissions arising from translation and adaptation or otherwise.

**Open access** This is an open access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited, appropriate credit is given, any changes made indicated, and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/.

**ORCID iDs**
Scott R Garrison http://orcid.org/0000-0002-7024-2158
Lee Green http://orcid.org/0000-0002-1789-7366
Dee Mangin http://orcid.org/0000-0003-2149-9376
Sabrina T Wong http://orcid.org/0000-0002-9619-9012
Rita McCracken http://orcid.org/0000-0002-2962-0364

**REFERENCES**

1. Veerman DP, Imholz BP, Wieling W, et al. Circadian profile of systemic hemodynamics. Hypertension 1995;26:55–9.
2. Clement DL, De Buyzere ML, De Bacquer DA, et al. Prognostic value of ambulatory blood-pressure recordings in patients with treated hypertension. N Engl J Med 2003;348:2407–15.
3. Verdecchia P, Porcellati C, Schillaci G, et al. Ambulatory blood pressure: an independent predictor of prognosis in essential hypertension. Hypertension 1994;24:793–801.
4. Ben-Dov IZ, Kard JD, Ben-Ishay D, et al. Predictors of all-cause mortality in clinical ambulatory monitoring: unique aspects of blood pressure during sleep. Hypertension 2007;49:1235–41.
5. Fagard RH, Celis H, Thijs L, et al. Daytime and nighttime blood pressure as predictors of death and cause-specific cardiovascular events in hypertension. Hypertension 2008;51:55–61.
6. Hermida RC, Ayala DE, Fernández JR, et al. Administration-time differences in effects of hypertension medications on ambulatory blood pressure regulation. Chronobiol Int 2013;30:280–314.
7. Hermida RC, Ayala DE, Mojon A, et al. Influence of circadian time of hypertension treatment on cardiovascular risk: results of the MAPEC study. Chronobiol Int 2010;27:1629–51.
8. Nerenberg KA, Zimke KB, Leung AA, et al. Hypertension Canada’s 2018 guidelines for diagnosis, risk assessment, prevention, and treatment of hypertension in adults and children. Can J Cardiol 2018;34:506–25.
9. Unger T, Borghi C, Charchar F, et al. 2020 International Society of Hypertension global hypertension practice guidelines. Hypertension 2020;75:1334–57.
10. Whelton PK, Carey RM, Aronow WS, et al. 2017 ACC/AHA/ABC/ACP/AAPA/AGS/ASH/ASPC/NMA/PCNA guideline for the prevention, detection, evaluation, and management of high blood pressure in adults: a report of the American College of Cardiology/American heart association Task force on clinical practice guidelines. Hypertension 2018;71:e113–115.
11. NICE Guidelines. Hypertension in adults: diagnosis and management. Available: https://www.nice.org.uk/guidance/ng1 [Accessed 13 Dec 2021].
12. Burnier M, Kreutz R, Narkiewicz K, et al. Circadian variations in blood pressure and their implications for the administration of antihypertensive drugs: is dosing in the evening better than in the morning? J Hypertens 2020;38:1396–406.
13. Turgeon RD, Althouse AD, Cohen JB, et al. Lowering Nighttime Blood Pressure With Bedtime Dosing of Antihypertensive Medications: Controversies in Hypertension - Con Side of the Argument. Hypertension 2021;78:871–8.
14. Hermida RC, Ayala DE, Fernández JR, et al. Comparison of the efficacy of morning versus evening administration of telmisartan in essential hypertension. Hypertension 2007;50:715–22.
15. Hermida RC, Ayala DE, Fontao MJ, et al. Administration-time-dependent effects of spirapril on ambulatory blood pressure in uncomplicated essential hypertension. Chronobiol Int 2010;27:560–74.
16. Hermida RC, Ayala DE, Fontao MJ, et al. Chronotherapy with valsartan/amlopidine fixed combination: improved blood pressure control of essential hypertension with bedtime dosing. Chronobiol Int 2010;27:1287–303.
17. Hermida RC, Ayala DE, Mojon A, et al. Reduction of morning blood pressure surge after treatment with nifedipine GITS at bedtime, but not upon awakening, in essential hypertension. Blood Press Monit 2009;14:152–9.
18. Hermida RC, Ayala DE, Mojon A, et al. Comparison of the effects on ambulatory blood pressure of awakening versus bedtime administration of torasemide in essential hypertension. Chronobiol Int 2008;25:950–70.
19. Hermida RC, Ayala DE, Mojon A, et al. Chronotherapy with nifedipine GITS in hypertensive patients: improved efficacy and safety with bedtime dosing. Am J Hypertens 2008;21:948–54.
20. Hermida RC, Ayala DE, Mojon A, et al. Ambulatory blood pressure control with bedtime aspirin administration in subjects with prehypertension. Am J Hypertens 2009;22:896–903.
21. Hermida RC, Ayala DE, Chayan L, et al. Administration-time-dependent effects of olmesartan on the ambulatory blood pressure of essential hypertension patients. Chronobiol Int 2009;26:61–79.
22. Hermida RC, Crespo JJ, Dominguez-Sardiña M, et al. Bedtime hypertension treatment improves cardiovascular risk reduction: the Hygia chronotherapy trial. Eur Heart J 2020;41:4565–76.
23. Kreutz R, Kjeldsen SE, Burnier M, et al. Disregard the reported data from the HYGIA project: blood pressure medication not to be routinely dosed at bedtime. J Hypertens 2020;38:2144–5.
24. Brunstrom M, Kjeldsen SE, Kreutz R, et al. Missing verification of source data in hypertension research: the HYGIA project in perspective. Hypertension 2021;78:555–8.
25. Carlsberg B, Brunstrom B. Is bedtime the best time of day? International Society of hypertension March 2020 Newsletter. Available: https://sh-world.com/data/uploads/2003-1.pdf?page=19 [Accessed 17 Dec 2021].
26. Chan A-W, Tetzlaff JM, Gotzsche PC, et al. SPIRIT 2013 explanation and elaboration: guidance for protocols of clinical trials. BMJ 2013;346:e7586.
27. Hennson L, Heiner T, Dahlböf B. Prospective randomized open blinded end-point (probe) study. A novel design for intervention trials. prospective randomized open blinded end-point. Blood Press 1992;1:113–9.
28. Grieshaber MC, Flammer J. Blood flow in glaucoma. Curr Opin Ophthalmol 2005;16:79–83.
29. Hayreh SS. Role of nocturnal arterial hypertension in the development of ocular manifestations of systemic arterial hypertension. Curr Opin Ophthalmol 1999;10:474–82.
30. Hayreh SS, Zimmerman MB, Podhajsky P, et al. Nocturnal arterial hypertension and its role in optic nerve head and ocular ischemic syndromes. Am J Ophthalmol 1994;117:603–8.
31. Harris PA, Taylor R, Thielke R, et al. Research electronic data capture (REDCap)--a metadata-driven methodology and workflow process for providing translational research informatics support. J Biomed Inform 2009;42:377–81.
32. Okutota RA, Hill MD. Coding of stroke and stroke risk factors using International classification of diseases, revisions 9 and 10. Stroke 2005;36:1776–81.
33 Quan H, Sundararajan V, Halfon P, et al. Coding algorithms for defining comorbidities in ICD-9-CM and ICD-10 administrative data. *Med Care* 2005;43:1130–9.

34 Tonelli M, Wiebe N, Fortin M, et al. Methods for identifying 30 chronic conditions: application to administrative data. *BMC Med Inform Decis Mak* 2016;15:31.

35 WebMD. Diuretics (water pills) for high blood pressure. Available: https://www.webmd.com/hypertension-high-blood-pressure/guide/diuretic-treatment-high-blood-pressure [Accessed 1 Jul 2021].

36 Heart and Stroke Foundation of Canada. Diuretics. Available: https://www.heartandstroke.ca/heart-disease/treatments/medications/diuretics [Accessed 1 Jul 2021].

37 Maasland L, van Oostenbrugge RJ, Franke CF, et al. Patients enrolled in large randomized clinical trials of antiplatelet treatment for prevention after transient ischemic attack or ischemic stroke are not representative of patients in clinical practice. *Stroke* 2009;40:2662–8.

38 Masoudi FA, Havranek EP, Wolfe P, et al. Most hospitalized older persons do not meet the enrollment criteria for clinical trials in heart failure. *Am Heart J* 2003;146:250–7.

39 Ruokoniemi P, Sund R, Arffman M, et al. Are statin trials in diabetes representative of real-world diabetes care: a population-based study on statin initiators in Finland. *BMJ Open* 2014;4:e005402.

40 Travers J, Marsh S, Williams M, et al. External validity of randomised controlled trials in asthma: to whom do the results of the trials apply? *Thorax* 2007;62:219–23.

41 Travers J, Marsh S, Caldwell B, et al. External validity of randomized controlled trials in COPD. *Respir Med* 2007;101:1313–20.

42 Mueller PS, Montori VM, Bassler D, et al. Ethical issues in stopping randomized trials early because of apparent benefit. *Ann Intern Med* 2007;146:878–81.

43 Rorie DA, Rogers A, Mackenzie IS, et al. Methods of a large prospective, randomised, open-label, blinded end-point study comparing morning versus evening dosing in hypertensive patients: the treatment in morning versus evening (TIME) study. *BMJ Open* 2016;6:e010313.

44 Garrison SR. BedMed-Frail: does the potential benefit of bedtime antihypertensive prescribing extend to frail populations? National Institutes of health clinical trials registry. Available: https://www.clinicaltrials.gov/ct2/show/NCT04054648 [Accessed 12 Jan 2021].