Screening for Hypertension in the INpatient Environment (SHINE): a protocol for a prospective study of diagnostic accuracy among adult hospital patients

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ABSTRACT

Introduction A significant percentage of patients admitted to hospital have undiagnosed hypertension. However, present hypertension guidelines in the UK, Europe and USA do not define a blood pressure threshold at which hospital inpatients should be considered at risk of hypertension, outside of the emergency setting. The objective of this study is to identify the optimal in-hospital mean blood pressure threshold, above which patients should receive postdischarge blood pressure assessment in the community.

Methods and analysis Screening for Hypertension in the INpatient Environment is a prospective diagnostic accuracy study. Patients admitted to hospital whose mean average daytime blood pressure after 24 hours or longer meets the study eligibility threshold for mean daytime blood pressure (≥120/70 mm Hg) and who have no prior diagnosis of, or medication for hypertension will be eligible. At 8 weeks postdischarge, recruited participants will wear an ambulatory blood pressure monitor for 24 hours. Mean daytime ambulatory blood pressure will be calculated to assess for the presence or absence of hypertension. Diagnostic performance of in-hospital blood pressure will be assessed by constructing receiver operator characteristic curves from participants’ in-hospital mean systolic and mean diastolic blood pressure (index test) versus diagnosis of hypertension determined by mean daytime ambulatory blood pressure (reference test).

Ethics and dissemination Ethical approval has been provided by the National Health Service Health Research Authority South Central—Oxford B Research Ethics Committee (19/SC/0026). Findings will be disseminated through national and international conferences, peer-reviewed journals and social media.

INTRODUCTION

Hypertension is the leading risk factor for death globally with 12.8% of annual mortality attributable to hypertension. A significant percentage of patients admitted to hospital have undiagnosed hypertension. However, unless a person has severely elevated blood pressure (BP), the clinical diagnosis of hypertension cannot be made in the in-hospital setting based on the UK, European or American guidelines. Indeed, increased BP values in hospital are frequently dismissed and additional potential reasons for this include clinicians attributing increased blood pressure values to anxiety, pain or white coat hypertension. However, the evidence suggests that patients with elevated BP recordings in hospital frequently remain hypertensive in the community. Despite this, referral for community follow-up of these patients to determine the presence or absence of persistent hypertension is poor.

Untreated hypertension is associated with a progressive increase in BP that can become treatment resistant. Therefore, hospital detection and subsequent management of hypertension offer an important intervention opportunity to address this major cause of morbidity and mortality.
In 2015, Oxford University Hospitals NHS Foundation Trust (OUHFT) introduced the NIHR-funded System for Electronic Notification and Documentation (SEND). SEND is a tailored software application, which links hospital bedside monitoring devices, including BP monitors, with a tablet computer for the manual recording of vital-sign observations of patients. Since this implementation, clinicians and researchers have been able to access and analyse patient observation data at a population as well as individual level. SEND has been used to record the vital-sign observations of more than 200,000 patients so far and to link each patient’s observations with their electronic patient record (EPR). This has paved the way for real time and automated recognition of patients whose observations indicate they may be at increased risk of undiagnosed hypertension.

The objective of this study is to identify the optimal in-hospital mean BP threshold, above which patients should receive postdischarge BP assessment in the community.

**METHODS AND ANALYSIS**

We prepared this protocol following the Standards for Reporting of Diagnostic Accuracy Studies (STARD) guidelines.

**Patient and public involvement**

The design of this study has been reviewed and supported by the patient and public representatives who have provided input on the research questions, data analysis, informational material for public engagement and the burden of the follow-up diagnostic testing from a patient’s perspective. Their input led to additional steps in data analysis, refining of the informational material and modification of the follow-up plan to reduce the patient burden.

**Study design**

This is a UK single-centre, prospective cohort diagnostic accuracy study. Recruitment commenced in November 2019 and active follow-up will continue until January 2022.

**Study setting**

Recruitment for this study will take place in OUHFT, UK. Participants will be followed up in a community research clinic at 8 weeks (±4) following discharge from hospital to receive reference BP testing. Participant timeline is shown in figure 1.

**Study population**

We will recruit participants aged between 18 and 80 years, who have at least two BP measurements taken during daytime hours defined as 07:00:00 to 21:59:59 hours and at least one BP measurement taken during night-time hours (22:00:00 to 06:59:59 hours). To be eligible, a patient’s cumulative mean daytime systolic BP must be between 120 mm Hg and 179 mm Hg and cumulative mean daytime diastolic BP ≤109 mm Hg for any given 24 hours interval since their admission to hospital following the first 24 hours of their admission (index test)—see the TEST METHODS section for more details. The mean daytime BP calculation must be based on a minimum of two blood pressure measurements made during daytime hours. No lower eligibility threshold was set for the diastolic blood pressure in order that patients with isolated systolic hypertension would not be excluded. Patients with a diagnostic code for hypertension in their medical record, as classified by the 10th revision of the International Statistical Classification of Diseases and Related Health Problems (ICD-10) or BP lowering medication prescribed in their medication record will be excluded. Additional exclusion criteria include current, intended or recent pregnancy, cause for index admission being associated with end-organ damage related to severe hypertension (including but not limited to heart failure, myocardial infarction, stroke, hypertensive encephalopathy) and estimatedglomerular filtration rate of <30 mL/min. Full inclusion and exclusion criteria are provided in table 1.

**Screening procedure**

The identification of patients whose BP exceeds the index test threshold will be automated using live, computerised algorithms. Blood pressure measurements in the OUHFT are made using automated oscillometric devices and the BP monitor readings (systolic and diastolic) are entered manually using the SEND computer tablet before automated transfer to the EPR by the SEND software app. The identification of patients whose BP measurements meet eligibility criteria will be computerised and automated. These automated systems for BP documentation and evaluating study inclusion criteria are intended to ensure that the measurement of the index test is objective and thus the clinical staff who perform the BP measurements can be considered blinded in the assessment of the index test. As this study is prospective in design, they will not be aware of the result of the reference standard.

**Participant sampling**

Sampling will be stratified to ensure that there is representation of patients with a full range of in-hospital BP profiles (according to predefined mean daytime systolic blood pressure, table 2). Patients who are considered eligible according to the screening algorithm will be approached by a clinical researcher who will undertake a further screen of patient eligibility. Patients who are in significant or uncontrolled pain will not be recruited unless and until, their pain improves. Pain scales will be collected at baseline for all recruited participants. Where there is insufficient study resource to enrol all recruitable patients, priority will be given to the strata with the least recruited patients. Recruitment will continue until the target recruitment number within

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**STARD** Standards for Reporting of Diagnostic Accuracy Studies

**BMJ Open** Open access journal

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each of the BP intervals is achieved. The prevalence of each systolic BP interval among the hospital population will be recorded within a screening log during the study period.

**Data collection**

Baseline data collection will be performed at the point of recruitment during the hospital admission, after the participant has provided informed, written consent. Custom-made baseline case report forms will be completed in the study data capture system for each participant by an appropriately qualified clinical researcher. The study data capture system will be electronic and accessible online, using a combination of data extracted from the EPR (e.g., diagnostic codes, current medications, serum haematology and biochemistry results and ECG) and participant self-report (e.g., ethnicity, family history, previous prescribed medications for hypertension). Any extreme values entered into the electronic data capture system will be inspected. A data monitoring committee has not been convened since there is no concealed allocation to treatment in this cohort study. Frequency and procedures for auditing study conduct are determined by the study sponsor (University of Oxford) based on a proportionate approach to risks.
The index test will be the first occurrence during the index admission of a cumulative mean daytime in-hospital BP reading ≥135 mm Hg systolic or ≥85 mm Hg diastolic. The systolic BP eligibility threshold has been defined as 10 mm Hg below the index test threshold to understand the frequency of true and false negatives in order to calculate sensitivity and specificity of the diagnostic test.

BP measurements from the point of admission to the time of assessment by the algorithm (every 24 hours at midnight until day 10 of the admission or discharge, whichever occurred first) will be averaged to reduce the chances of a person becoming eligible by chance when compared with the alternative, of assessing a person’s eligibility based on their mean day time BP for any 24 hours interval.

Thus, for each patient at each time of assessment, we will compute the cumulative mean BP reading using all the available daytime (7:00:00-21:59:59 hours) BP readings. The cumulative mean is defined here to be the mean of the mean daytime BP values, the latter being calculated for each of the days between the point of admission and the time of assessment. As the first occurrence of the test being passed is a necessary and sufficient condition, a patient will be recruited into the study even if their cumulative mean at discharge is below the index test thresholds (≥135 mm Hg systolic or ≥85 mm Hg diastolic).

A day time definition was chosen as this study is being performed in a UK healthcare setting, where at present, night-time measurements are not used for the diagnosis of hypertension. Hypertension. The index test has been defined according to the UK National Institute for Health and Care Excellence (NICE) guideline definition of daytime hypertension.

Additionally, a subgroup of participants will receive a research standard BP assessment in hospital, performed according to the standard recommended by the UK NICE guidelines. The BP will be measured using a validated and calibrated oscillometric device by a trained clinical researcher. BP will be measured with the patient seated and the arm outstretched and supported. BP will be measured in both arms and if a between-arm difference of ≥15 mg Hg is found, both measurements will be repeated. Where a difference persists, the arm with the highest measurement will be used for subsequent measurements. Two measurements will be taken in the selected arm; if the second measurement is substantially different from the first then a third measurement will be taken and the lowest of the second and third measurements will be recorded.

### Table 1 Full inclusion and exclusion criteria

<table>
<thead>
<tr>
<th>Inclusion criteria</th>
<th>Exclusion criteria</th>
</tr>
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<tbody>
<tr>
<td>Aged 18–80</td>
<td>Pre-existing diagnosis of hypertension or attending hospital with acute end-organ damage related to severe, undiagnosed hypertension.</td>
</tr>
<tr>
<td>Admitted to hospital for an acute or elective medical or surgical condition.</td>
<td>Presence of atrial fibrillation or other pulse rate irregularity, which means ABPM is not appropriate.</td>
</tr>
<tr>
<td>Have at least three in-hospital blood pressure recordings with two performed during day time hours and 1 performed during night-time hours, over a minimum of 24 hours for the index admission.</td>
<td>Currently pregnant, within 3 months postpartum or planning pregnancy during study period.</td>
</tr>
<tr>
<td>Identified by the blood pressure algorithm for any 24 hours interval to have a mean blood pressure which meets the eligibility thresholds.</td>
<td>Receiving treatments which might be used for the management of hypertension, for example, beta blockers for migraine, ACE inhibitors for renal disease.</td>
</tr>
<tr>
<td>Registered with a general practitioner.</td>
<td>Diagnosed with terminal illness or cognitive impairment.</td>
</tr>
</tbody>
</table>

Diagnosed with AKI on index admission or eGFR <30 mL/min. Cause for hypertension being toxicology, medical or recreational for example, amphetamines and their derivatives or alcohol withdrawal syndrome.

Postdischarge destination being another hospital or prison. Receiving concomitant chemotherapy. Already recruited to a separate hypertension study.

ABPM, ambulatory blood pressure monitoring; AKI, acute kidney injury; eGFR, estimated glomerular infiltration rate.

### Table 2 Systolic blood pressure intervals against which recruitment will be stratified

<table>
<thead>
<tr>
<th>Systolic blood pressure interval</th>
<th>Target recruitment no</th>
</tr>
</thead>
<tbody>
<tr>
<td>120–129</td>
<td>Approximately equal representation of participants recruited to each of these systolic blood pressure intervals</td>
</tr>
<tr>
<td>130–139</td>
<td></td>
</tr>
<tr>
<td>140–149</td>
<td></td>
</tr>
<tr>
<td>150–159</td>
<td></td>
</tr>
<tr>
<td>160–179</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>200</td>
</tr>
</tbody>
</table>
Reference standard diagnostic test

The gold-standard diagnostic test for the presence of hypertension will be a daytime 24-hour ambulatory BP threshold of ≥135 mm Hg systolic or ≥85 mm Hg diastolic, which is the current UK NICE diagnostic threshold for ambulatory blood pressure monitoring (ABPM).5

ABPM will be performed with each participant at 8 weeks (±4) following discharge from hospital using a validated automated Mobil-O-Graph ABPM (IEM, Stolberg, Germany), calibrated to manufacturer standards. Should the participant have acquired a diagnosis of hypertension and initiated treatment for hypertension following hospital discharge but before they receive ABPM, they will not be eligible to proceed with ABPM. In this instance, a participant will be regarded as being reference test positive, having acquired a clinical diagnosis of hypertension. The monitor will be programmed to measure BP twice per hour during waking hours for (07:00–22:00 hours), and once per hour during the sleeping hours (22:00–07:00 hours) for a period of 24 hours, according to the British and Irish Hypertension Society (BIHS) Standard Operating Procedure for performing ABPM.26 Adjustments will be made, for example, where a participant will be working a shift pattern during the period of wear. All participants will be asked to log their sleeping and waking times in an ABPM diary and these will be used to analyse the ABPM results. According to the BIHS standard operating procedure for ABPM, a minimum of 14 waking time ABPM measurements must be successful to calculate the mean daytime BP. Where fewer than 14 of the waking time ABPM measurements are successful, the participant will be asked to wear the monitor for a further period in order to collect adequate data. In the instance of two episodes of ABPM not collecting sufficient data, the participant and their registered general practitioner (GP) will be informed along with recommendation that the participant arrange an appointment with their GP for further evaluation of their BP. The participant will continue to be followed up remotely through the online data capture system for up to 52 weeks from recruitment (figure 1). Links to the questionnaires will be sent to participants by email and we will collect data regarding BP-related health outcomes, the prescription of antihypertensive medications and repeat blood pressure measurements. At 10 years, data will be obtained from the Office of National Statistics and Hospital Episode Statistics Databases regarding mortality and cardiovascular health-related outcomes.

The fitting and removal of the ABPM will be performed by a qualified clinical research nurse who is appropriately trained in the National Institute for Health’s Good Clinical Practice and to perform ABPM. At the time of removal of the monitor, the electronic ABPM report will be downloaded to the study data capture system and interpreted by the research nurse. As the reference test procedure is objective through the use of an oscillometric, automated ABPM, the research nurse will not be blinded to the result of the index BP test.

The same subgroup of 50 participants who received a standardised research BP assessment in hospital will receive a standardised research clinic blood pressure on ABPM fitting. The procedure for measuring and recording the BP will be exactly the same as described for the hospital setting, according to that recommended by the UK NICE guidelines.27

STATISTICAL METHODS

Sample size

Owing to the novel design of this study, estimates of expected prevalence of hypertension at follow-up have been used to calculate sample size. From a retrospective analysis of in-hospital OUHFT BP data, we estimate that 3000 patients per annum will be eligible for study inclusion (Mahdi, A, 2019; Prevalence of hypertension and undiagnosed hypertension in a large inpatient population). Previous studies have used an in-hospital BP eligibility threshold (140/90 mm Hg) and revealed that approximately half of the patients with a BP above the threshold have hypertension at follow-up.28 29 Although this study uses a lower eligibility BP threshold of 120/70 mm Hg; we estimate half of the patients will have hypertension at follow up owing to the stratification used for recruitment. Therefore, a sample size of 196 participants has been estimated to permit detection of a 50% rate of hypertension at follow-up, with a 95% confidence interval width of approximately ±7%. Screening logs will be recorded for both the algorithm and the manual screening process to understand the prevalence of patients falling in each of the in-hospital BP intervals against which recruitment will be stratified.

Baseline characteristics

Clinical and demographic characteristics of the study population will be reported, including age, sex, presenting complaint, comorbidity, current treatments (including oral steroidal and non-steroidal anti-inflammatory medications), pain score (using an 11-point Likert scale where 0=no pain and 10=maximum possible pain) for the preceding 24 hours to recruitment and the Six-Item State Anxiety Scale.30 The number of participants who complete each stage of the protocol will be reported, including those who complete successful ABPM, with reasons for failed testing where appropriate. Recruitment and study completion figures will be reported using a participant flow diagram. Interpretation of the ABPM result will be performed by the researchers who are blind to the participant’s index BP test.

Index test performance

Diagnostic performance of in-hospital BP will be assessed by constructing receiver operator characteristic (ROC) curves from participants’ in-hospital mean systolic or mean diastolic BP (index test) versus diagnosis of hypertension determined by mean daytime ambulatory systolic and diastolic BP (reference test). At first instance, a complete-case analysis will be performed by excluding...
participants without an observed test result. At the second instance, for participants with missing ABPM values, the data will be imputed. The two approaches will be compared and the number of cases with missing values will be reported, together with detailed description of the strategy for handling the missing data. The area under the ROC curve will be calculated to estimate the discriminatory power of the diagnostic test. Potential diagnostic BP thresholds will then be identified from the ROC curve. The diagnostic ability of each potential cut-off threshold will be evaluated using two-by-two tables to compare sensitivity and specificity of each potential threshold. Confidence intervals will be reported for each value. Reproducibility of the index test and variability in diagnostic accuracy will be analysed using subsequent available periods of the hospital admission in which participants’ mean in-hospital BP values were at or above the recruitment threshold.

**Subgroup analyses**

Logistic regression analyses will be performed to estimate whether the diagnostic performance of in-hospital mean BP is influenced by participant baseline characteristics, including age, sex, body mass index and comorbidity, pain, anxiety and in-hospital prescriptions such as oral steroid medications and non-steroidal anti-inflammatory medications. Further subgroup analyses will include whether the index admission was scheduled or unscheduled and whether the admitting specialty was medical or surgical.

**Additional analyses**

Once the optimal thresholds for the in-hospital screening for hypertension have been identified, diagnostic accuracy of the thresholds at successive 24 hours intervals of patient admission will be estimated. While the reference test for this study is based on the current UK diagnostic criteria for the diagnosis of hypertension, which concerns daytime BP measurements only, there is increasing evidence in the literature to support the association between nocturnal hypertension and cardiovascular risk. Therefore, additional analyses will be performed incorporating night-time measurements and using the European and American diagnostic thresholds for hypertension as shown in table 3.

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**Table 3** European Society of Cardiology/European Society of Hypertension and American College of Cardiology diagnostic thresholds for hypertension

<table>
<thead>
<tr>
<th>ABPM diagnostic thresholds for hypertension</th>
<th>European Society of Cardiology/European Society of Hypertension</th>
<th>American College of Cardiology</th>
</tr>
</thead>
<tbody>
<tr>
<td>Systolic</td>
<td>Diastolic</td>
<td>Systolic</td>
</tr>
<tr>
<td>24hours</td>
<td>130</td>
<td>80</td>
</tr>
<tr>
<td>Night</td>
<td>120</td>
<td>70</td>
</tr>
<tr>
<td>Day</td>
<td>135</td>
<td>85</td>
</tr>
</tbody>
</table>

ABPM, Ambulatory Blood Pressure Monitoring.

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**ETHICS AND DISSEMINATION**

Any planned protocol modifications will receive further sponsor and ethical review and require approval prior to implementation. Any approved modifications will be communicated in the ISRCTN Registry where the study is registered (Identification number: ISRCTN80586284). All participants will be required to give informed, written consent and will be advised that they can withdraw from the study at any time and without giving reason. Consent will be obtained by a trained clinical researcher.

We do not anticipate any harms from the use of ABPM, which is already a standard procedure for the diagnosis of hypertension in UK primary care, and therefore, this study does not carry any significant risk to participants. The measurement of ambulatory BP among the participants of this study is in addition to usual care and may be considered beneficial if we detect the presence of undiagnosed hypertension earlier than may have otherwise been the case. If at any point a participant is found to have severely raised blood pressure, their responsible hospital care team or GP will be informed, following study standard operating procedures, according to whether severely elevated BP was observed in the hospital or community, respectively. Any adverse events, which arise as a result of performing ABPM, will be reported. The participant and their GP will both be informed of the ABPM result and any decision to initiate treatment will be made between the patient and their GP in line with standard care.

Data will be analysed by researchers based in the University of Oxford (Nuffield Department of Clinical Neurosciences, Nuffield Department of Primary Care Health Sciences and Institute of Biomedical Engineering).

**Dissemination**

Findings from this study will be disseminated through peer-reviewed journals, conferences, public engagement activities and online. Participants will be able to access results online.

**Discussion**

The Screening for Hypertension in the INpatient Environment study will be the first study to our knowledge, to evaluate the diagnostic accuracy of in-hospital BP measurements on a continuous scale and therefore identify the optimal BP cut-off values for in-hospital...
hypertension screening. A recent systematic review indicates this will be the first study to use real-time algorithms to screen hospital inpatients’ electronic BP data. The automated detection and subsequent treatment of previously undiagnosed hypertension has the potential to prevent the development of serious hypertension-related diseases.

The study is limited by uncertainty in the expected prevalence of hypertension at follow-up, which has been used to estimate the required sample size.

**REFERENCES**

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