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Study protocol for Smartphone Monitoring for Atrial fibrillation in Real-Time in India (SMART-India): a community-based screening and referral programme

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ABSTRACT

Introduction Atrial fibrillation (AF), the world’s most common arrhythmia, often goes undetected and untreated in low-resource communities, including India, where AF epidemiology is undefined. AF is an important risk factor for stroke, which plagues an estimated 1.6 million Indians annually. As such, early detection of AF and management of high-risk patients is critically important to decrease stroke burden in individuals with AF. This study aims to describe the epidemiology of AF in Anand District, Gujarat, India, characterise the clinical profile of individuals who are diagnosed with AF and determine the performance of two mobile technologies for community-based AF screening.

Methods This observational study builds on findings from a previous feasibility study and leverages two novel technologies as well as an existing community health programme to perform door-to-door AF screening for 2000 people from 60 villages of Anand District, Gujarat, India using local health workers. A single-lead ECG and a pulse-based application is used to screen each individual for AF three times over a period of 5 days. Participants with suspected arrhythmias are followed up by study cardiologist who makes final diagnoses. Participants diagnosed with AF are initiated on treatment based on current anticoagulation guidelines and clinical reasoning.

Analytical plan Age-stratified and sex-stratified prevalence of AF in the Anand District will be calculated for sample and estimated for Anand distribution using survey design weights. Sociodemographic and clinical factors associated with AF will be evaluated using multivariable regression methods. Performance of each mobile technology in detecting AF will be evaluated using 12-lead ECG interpretation as the gold standard.

Ethics and dissemination This protocol was approved separately by the Institutional Review Board of University of Massachusetts Medical School and the Human Research Ethics Committee at Charutar Arogya Mandal. The findings of this study will be disseminated through peer-reviewed journals and scientific conferences.

Strengths and limitations of this study

► The Smartphone Monitoring for Atrial fibrillation in Real-Time in India study represents an innovative screening and referral paradigm for persons living in low-resource settings who may have an undetected chronic disease.
► This study highlights the importance of conducting a feasibility study and engaging local community partners prior to undertaking more broad-based population screening efforts.
► Serial screening and rigorous clinical follow-ups, which include continuous monitoring using a Holter device, increase the chances of identifying paroxysmal atrial fibrillation (AF) events. By contrast, most AF screening programmes described in the literature rely on a single-point screening.
► Our efforts to characterise the clinical profile of participants diagnosed with AF are limited by the cross-sectional design of our study which prohibit us from drawing casual inferences.

BACKGROUND

Clinical and public health significance

Although atrial fibrillation (AF) is the world’s most common serious heart rhythm problem, it often goes undetected and untreated, particularly in low-resource settings like India.1–5 As such, little is known about the epidemiology of AF in India.1–3 5 However, it is estimated that five million Indians have AF, based on ageing of the Indian population and limited data showing that the Indian population is increasingly affected by AF risk factors, including hypertension, diabetes mellitus and coronary heart disease.6 The importance of AF to Indian public health is underscored...
by an ongoing stroke epidemic with an estimated 1.6 million Indians experiencing a stroke in 2015. AF is a major risk factor for stroke conferring a fivefold higher risk among affected individuals compared with those free from the arrhythmia, suggesting that AF has become a major public health concern in India.6–10

Public health programmes for the routine surveillance of AF or other risk factors for stroke are sparse in India. A small proportion of Indian residents have access to routine cardiovascular care or prompt stroke treatment, and this subgroup is disproportionately clustered in urban and metropolitan cities. In light of the limited access to, and out-of-pocket costs associated with, stroke treatment, AF screening and treatment represent an appealing method for primary stroke prevention in rural India.7 Targeted AF screening has been employed successfully in high-income countries, such as the UK and Australia,11–14 but there are no major systematic efforts to screen for AF in low-income and middle-income countries. Furthermore, oral anticoagulation to prevent stroke is an effective therapy for patients with AF, but solutions are needed to assist those with limited means to pay for this treatment.5 15–17

Another major barrier to AF screening has been the resource intensiveness of performing conventional 12-lead ECG to diagnose AF.18 For these and other reasons, the importance of developing new, affordable, mobile devices for AF screening that could be deployed in India was emphasised by an National Heart, Lung, and Blood Institution Expert panel and at the 2008 Hyderabad workshop.3

In this manuscript, we present an innovative community-based study called Smartphone Monitoring for Atrial fibrillation in Real-Time—India (SMART-India). Our approach deploys two novel smart device-based arrhythmia analysis technologies for AF screening using an existing network of community-health workers in rural Western India, medical professionals and provincial health leaders, and builds on a foundational feasibility assessment conducted in Gujarat, India. The SMART-India study is supported by our interdisciplinary Indo-US research team that has an ongoing collaboration between investigators with a proven track record of conducting community-based technology research in India.19

AF screening technology

Single-lead ECG device

Kardia’s Food and Drug Administration (FDA) approved AliveCor device records a single-lead ECG signal from the user. Participants place two or three fingers of their right hand over the standard camera and first finger or second finger over the standard camera and lamp. The flashlight illuminates the finger and the pulse waveform is recorded for up to 2 min, with 30 s being needed in most cases to complete a rhythm analysis. The ANAND algorithm performs two calculations to approximate a person’s cardiac rhythm: (1) root mean square of successive difference of RR intervals to quantify variability between RR peaks and (2) Shannon entropy (or turning point ratio) to characterise the complexity of variability between two consecutive segments of 14 RR peaks.22 The results from this automated algorithm’s calculation on the waveform are classified as normal sinus rhythm, possible AF or indeterminate.21 The preceding prototype for ANAND application (app) was tested in a clinical environment and compared its performance against a manually adjudicated ECG reading demonstrating a sensitivity of 0.97 and a specificity of 0.94 for the detection of AF. This ANAND app is not commercially available and differs from the early prototype in two main aspects: a shorter duration of pulse recording and optimised threshold for RR variability and Shannon entropy based on data collected from our feasibility study.23

Feasibility study

In mid-2015, a feasibility study was undertaken in rural Western India to screen for AF using the two smartphone-based technologies: (1) the AliveCor device and (2) the ANAND app. Our primary objective for this study was to help identify the potential barriers and facilitators for conducting AF screening on a larger, more diverse and representative sample.

Our field staff for each village consisted of two undergraduate medical students who were trained in collecting field observation notes (online supplementary appendix A), a research coordinator, two social workers who were trained in administering questionnaires and using mobile technology to screen for AF and a local village health worker to help identify residents of the given village who meet the eligibility criteria. We screened a total of 354 participants aged 50 years and above, across six villages in Anand District in Gujarat. Each participant in this pilot study was screened for 2 min using both technologies for five consecutive days. Data was collected over a 6-week period.

Pulse-based ANAND application

Our team has developed a novel pulse-based approach for detecting arrhythmias that does not require additional hardware. This experimental Automated Novel Atrial fibrillation Noninvasive Detection (ANAND) application, named after our study site in India, builds on our previous USA-based algorithm to identify pulse complexity and irregularity.21 22 Participants are asked to hold the mobile phone in their hand, with their right first finger or second finger over the standard camera and lamp. The flash light illuminates the finger and the pulse waveform is recorded for up to 2 min, with 30 s being needed in most cases to complete a rhythm analysis. The ANAND algorithm performs two calculations to approximate a person’s cardiac rhythm: (1) root mean square of successive difference of RR intervals to quantify variability between RR peaks and (2) Shannon entropy (or turning point ratio) to characterise the complexity of variability between two consecutive segments of 14 RR peaks.22 The results from this automated algorithm’s calculation on the waveform are classified as normal sinus rhythm, possible AF or indeterminate.21 The preceding prototype for ANAND application (app) was tested in a clinical environment and compared its performance against a manually adjudicated ECG reading demonstrating a sensitivity of 0.97 and a specificity of 0.94 for the detection of AF. This ANAND app is not commercially available and differs from the early prototype in two main aspects: a shorter duration of pulse recording and optimised threshold for RR variability and Shannon entropy based on data collected from our feasibility study.23

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Key findings from field observation notes and in-depth interviews of the field staff are described in table 1. The sample prevalence of AF using the clinician’s adjudication of single-lead ECG recordings from the AliveCor device was found to be 5.1% (95% CI 2.7% to 8.7%).

Community-based approach

Our approach builds on a long-standing collaboration between the University of Massachusetts Medical School and Charutar Arogya Mandal (CAM), a tertiary care hospital and medical college in rural Western India. This collaboration, which we have called Research and Community Engagement Activities (RACE), is grounded in extensive interactions with the local Indian communities. As we have described elsewhere, the research priorities of our collaboration and the local Indian communities. As we have described elsewhere,

Table 1 Findings from the feasibility study and corresponding modifications in the SMART-India study

<table>
<thead>
<tr>
<th>Theme</th>
<th>Findings</th>
<th>Modification</th>
</tr>
</thead>
</table>
| Sampling strategy      | ► Recruiting an equal distribution of participants across the different age–sex stratum was difficult  
► Approaching households in a random fashion to recruit participants resulted in a bias towards selection of middle-aged female and older participants who were more likely to be present at home and available for screening | Use of census data to recruit an age-representative and sex-representative sample through stratified sampling to mitigate potential selection bias |
| Motion artefact        | ► Participants found it difficult to maintain steady hands while placing their fingers on the AliveCor case without additional support  
► Participants found 2 min screening period to be too long to maintain steady hands | ► Use of clipboards to stabilise participants’ hands  
► Shorter recording time: 60 s (AliveCor) and 90 s (ANAND) |
| Screening protocol     | ► A single time-point screening underestimated true AF prevalence (owing to paroxysmal AF)  
► Five screenings in a week were time consuming and led to participant attrition | Three screenings over the course of 5 days to account for paroxysmal events and prevent study attrition |
| Data management        | ► Research staff had difficulties uploading ECG tracings for adjudication in a timely manner  
► Questionnaires in the field could not be entered using smartphones and tablets due to poor internet connectivity | ► Development of a web platform to assist with sharing of de-identified ECG recordings  
► Use of an off-line smartphone-based application |

AF, atrial fibrillation; ANAND, Automated Novel Atrial fibrillation Noninvasive Detection; SMART-India, Smartphone Monitoring for Atrial fibrillation in Real-Time—India.

METHODS/DESIGN

Objectives

Box lists the primary objectives of the SMART-India study. These objectives are informed by the feasibility study, AF-screening technology and community-based approach.

Study design

Figure 1 provides an overview of the SMART-India study protocol from the perspective of prospective participants. The SMART-India study is a cross-sectional screening of 2000 adult men and women aged 45 years and older.
Box Study aims for Smartphone Monitoring for Atrial fibrillation in Real-Time—India

- Aim 1: Describe the epidemiology of atrial fibrillation (AF) in Anand District, Gujarat, India. Screen 2000 residents of Gujarat to describe the age-stratified and sex-stratified prevalence of AF.
- Aim 2: Characterise the profile of individuals who are diagnosed with AF. Compare clinical findings, quality of life and risk factors of participants who are diagnosed with AF with those who do not have AF.
- Aim 3: Determine the performance of two mobile technologies for community-based AF screening. Determine the sensitivity, specificity and discriminative ability of the AliveCor and Automated Novel Atrial fibrillation Non-invasive Detection's automated AF detection algorithm to the gold-standard 12-lead ECG as well as interpretation by a trained clinician.

from 60 villages participating in the SPARSH programme paired with a clinical follow-up of all participants with abnormal screenings. Each participant is screened three times over the course of 5 days to account for potential paroxysmal AF presentation.

Staff training

The SMART-India study is supported by 4 field workers, 1 clinical coordinator, 1 project supervisor and 50 village health workers. The field workers and clinical coordinator received 2 weeks of training in the use of smartphones and devices to administer questionnaires to surveyed individuals and perform screening for AF. Additionally, the five staff members worked with the investigators to develop standardised operating procedures for their tasks. Following training, the five team members collaborated with the study investigators to organise a 1-hour orientation for the 60 village health workers. Participants from 50 villages took part in this study, while volunteers from the other 10 villages had participated in the feasibility study described earlier. Training provided the village health workers with information about the study, their roles and introduction to the rest of the study team. During the orientation, village health workers indicated that navigating the village to identify participants listed in the roster would be the primary time constraint. Thus, we divided the four field workers into two teams and proceeded with simultaneous screenings in two separate villages to enhance efficiency. One of the Indian co-investigators on the team fulfilled the role of project supervisor.

Questionnaire translation and testing

We use standardised questionnaires to collect information about participants’ demographic characteristics, lifestyle risk factors, past medical history, family history, physical activity, medication, AF-related symptoms and healthcare usage at the time of the first screening (Table 2). Additionally, we administer an exit survey to ascertain participants’ feedback about the screening technology and a clinical follow-up survey to assess participants’ quality of life and cardiovascular-related factors. All questionnaires were translated to Gujarati and interpreted by several volunteers representing a wide range of literacy (ie, attending physician, healthcare worker, research coordinator, village health worker and community representative). The interpretations of questions and responses were back-translated and compared with the original questionnaire in English. Common themes for discordance between the original and back-translated forms were identified and used to modify the original questionnaire to accommodate the cultural context. This process was repeated until there was convergence between the two versions. All translations were done by certified professional translator services in Gujarat, India.

Data management

Figure 2 describes the data management strategy for SMART-India study. Based on our prior experience, we use the Magpi platform to administer questionnaire-based surveys to the participants. Magpi allows for off-line data collection in smartphones and tablets through a user-friendly app and uploads collected data to a secure server once the device connects to the Internet. The files from the

Figure 1 An overview of SMART-India study protocol for community-based screening of AF and referral for clinical follow-up in rural Western India. AF, atrial fibrillation; ANAND, Automated Novel Atrial fibrillation Noninvasive Detection; app, application; FDA, Food and Drug Administration; SMART-India, Smartphone Monitoring for Atrial fibrillation in Real-Time—India; SPARSH, Shree Krishna Hospital Programme for Advancement of Rural and Social Health.
Table 2  Data collected using standardised questionnaires during enrolment, final screening and follow-up time points for Smartphone Monitoring for Atrial fibrillation in Real-Time—India study

<table>
<thead>
<tr>
<th>Enrolment</th>
<th>Variables</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sociodemographic characteristics</strong></td>
<td>Age, sex, household size, education level, occupation, caste, religion, marital status</td>
</tr>
<tr>
<td><strong>Financial status</strong></td>
<td>Household income, perceived income, perceived expenditures</td>
</tr>
<tr>
<td><strong>Healthcare</strong></td>
<td>Healthcare use, healthcare seeking behaviour, healthcare expenditures</td>
</tr>
<tr>
<td><strong>Lifestyle</strong></td>
<td>Smoking, non-smoking tobacco, alcohol, expenditures on lifestyle activities</td>
</tr>
<tr>
<td><strong>Physical activity</strong></td>
<td>Mild, moderate and high levels of activity using International Physical Activity Questionnaire</td>
</tr>
<tr>
<td><strong>Cardiovascular symptoms</strong></td>
<td>Palpitations, irregular heartbeats, skipped heartbeats, dizziness</td>
</tr>
<tr>
<td><strong>Medical history</strong></td>
<td>Valvular heart disease, coronary heart disease, obstructive lung disease, myocardial infarction, hypertension, diabetes mellitus, hypercholesterolaemia, heart failure, stroke, transient ischaemic attack, systemic embolism, anaemia, internal bleeding (intracranial, gastrointestinal or other), chronic kidney disease, congenital heart disease</td>
</tr>
</tbody>
</table>

Exit

| Technology access | Cell phone ownership, cell phone with camera, smartphone use, internet usage, internet use to find health information |
| ANAND usability | Time-consuming, operational ease, understandability of results |

Follow-up

| Quality of life | Physical and Mental composite score using SF-12 |
| Need for assistance | Recreational activities, travelling, shopping, preparing or eating meals, day-to-day activities, medication management, financial management |

ANAND, Automated Novel Atrial fibrillation Non-invasive Detection; SF-12, Short Form-12 Survey.

AF screening apps (AliveCor and ANAND) are downloaded to a personal computer from the phones using iTunes. Results from the automated algorithm of AF screening apps and all abnormal AliveCor ECG tracings are uploaded to REDCap. Additionally, all data emerging from the clinical follow-up, except for questionnaire-based surveys, are uploaded to REDCap. REDCap is used because it allows for file uploads and can create reports that are helpful during the adjudication process.

Sample size calculations

Based on our calculations, a sample size of 1823 persons is required to estimate the prevalence of AF with 1% error assuming the a priori prevalence of AF to be 5% based
on our feasibility study. The resultant stratum size of 300 people can estimate the subpopulation prevalence of 5% with a 95% CI of 2.8% to 8.1%. An effective screening tool maximises sensitivity to avoid false-negative findings. Assuming a 5% prevalence of AF, a sample size of 2000 participants (5% AF rate, n=100) or more is necessary to obtain a satisfactory confidence band for 90% sensitivity for study aim 3 (table 3). Considering the extent of resources available, the study team decided to screen 2000 adult men and women 45 years and older to accomplish the aims outlined in box.

Community-based screening

Prescreening preparation

One week prior to screening, field workers for study team A and team B contact the respective village health workers to inform them of the screening plan. Both teams develop an age-stratified and sex-stratified recruitment roster for their respective villages based on the census provided by the SPARSH programme. In total, six strata are defined for the combination of age (40–55, 56–65 and 65+ years) and sex (male and female) categories. Teams print out recruitment and screening tracking forms (online supplementary appendix B) and charge the smartphones and portable powerbanks for use during the week.

Screening protocol

Recruitment and enrolment occurs on day 1 and day 2 of the screening process. The two study teams accompany their respective village health workers to enrol 21 willing participants in their respective villages on a daily basis. Each participant is screened using the AliveCor device for 1 min, with the AliveCor device placed on the clipboard (figure 3), followed by the ANAND app for 1 min 30 s. Results are recorded in the tracking form. If the participant is screened for either unclassified or unreadable heart rhythm on AliveCor, two more attempts are made to obtain either a normal or possible AF automated reading. Following this screening process, each participant completes the enrolment questionnaire that is entered directly into the Magpi database. Village health workers perform random blood glucose testing and measured the blood pressure of all participants. On day 3, both field teams attempt to screen all 42 participants using the AliveCor device and ANAND app. A successful follow-up, number of attempts required and whether or not the screening occurred at home is recorded in the tracking form. On day 4 and day 5, the third screening for AF is again split into 2 days.

If a participant is not available on day 3, field workers approach them on both days to perform a total of three screenings over 5 days. Thus all participants are screened 1 and 2 days apart, unless they are unavailable during their follow-up screening. After the final screening was completed, the field workers administer an exit survey, and data is entered directly into Magpi forms. The results of all screenings are recorded on paper and entered into REDCap along with the AliveCor ECG recordings for those who have an abnormal finding.

Postscreening activities

On each of the final 2 days (day 4 and day 5) of screening, field workers travel back to CAM to complete data entry and upload all abnormal (possible AF/unclassified) ECG files to REDCap. All AliveCor and ANAND files are downloaded from the phones using iTunes. All AliveCor portable document formats are renamed with patient IDs and separated into normal results versus AF/unclassified results. All AF/unclassified files are uploaded to REDCap and flagged for adjudication. The field workers create a weekly report to summarise recruitment and results of screening and distribute it with the team (online supplementary appendix C). All of the ANAND and AliveCor files are uploaded to a secure server to support ad hoc analyses for measuring the performance of the two

Table 3 95% CI for sensitivity and specificity by number of participants to justify the sample size of 2000 participants for the Smartphone Monitoring for Atrial Fibrillation in Real-Time in India (Smart-India study (assumes 5% prevalent atrial fibrillation))

<table>
<thead>
<tr>
<th>Number of participants</th>
<th>500</th>
<th>1000</th>
<th>2000</th>
<th>5000</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sensitivity (0.9)</td>
<td>0.69 to 0.97</td>
<td>0.78 to 0.97</td>
<td><strong>0.82 to 0.95</strong></td>
<td>0.86 to 0.93</td>
</tr>
<tr>
<td>Specificity (0.9)</td>
<td>0.87 to 0.93</td>
<td>0.88 to 0.92</td>
<td><strong>0.89 to 0.91</strong></td>
<td>0.89 to 0.91</td>
</tr>
</tbody>
</table>

screening methods. Ultimately all data are exported to STATA for statistical analyses (figure 2).

**Clinical follow-ups**

Our protocol for clinical follow-ups evolved over time based on feedback from the research team. The primary impetus behind the change in approach was higher than expected number of screenings that are classified as inconclusive and the resultant backlog of follow-up visits needed to understand these inconclusive readings. In order to facilitate prompt clinical visits for high-risk participants, we introduced a triaging step (Modifications for follow-up scheduling) after reviewing the clinical charts and final diagnosis for the first 50 participants that were followed up in clinic.

**Scheduling follow-ups**

The clinical coordinator schedules all follow-up visits with village health workers. One day prior to the visit, the clinical coordinator informs the cardiac and extension departments about the number of participants and which village health workers are coming. The clinical coordinator prints out the abnormal ECGs and paper clinical forms to prepare for the follow-up visit.

**Follow-up protocol**

The clinical coordinator meets the village health workers and participants at the hospital and escorts them to the cardiac centre. Once at the cardiac centre, the clinical coordinator conducts further AF screening using the AliveCor device and ANAND app using the same methodology as in the field. Subsequently, all participants receive a 12-lead ECG. The study cardiologist reviews all of the ECG recordings (field and clinical visit) and determines whether or not to pursue additional clinical work-up. While waiting for the cardiologist’s interpretation, the clinical coordinator administers the follow-up survey in Magpi, recorded vital information as measured by the nurse (height, weight, blood pressure, heart rate, O₂ saturation, temperature) and verified the participants’ medical history. The cardiologists’ clinical decision-making is described in figure 4. If the cardiologist interprets all ECG recordings of the participant as showing normal sinus rhythm, no additional studies are ordered. If the cardiologist diagnosed the participant as having definite AF in any of the ECG, either in any of the screening ECG or in-office ECG, additional laboratory investigations are performed to inform clinical decision-making about initiation of anticoagulation and management plan (figure 4). If AF was suspected on any of the screening ECG but a diagnosis of definite AF could not be made due to poor quality of ECG tracings from screening app, multiple atrial or ventricular premature complexes, or a normal sinus rhythm on the 12-lead ECG, the cardiologist orders 24-hour Holter monitoring to look for paroxysmal AF events or other abnormalities. Two-dimensional echocardiography is performed in all participants diagnosed with AF and in those having symptoms and/or ECG abnormalities that warrants for an echocardiographic examination. Decision to initiate anticoagulation is taken based on Congestive heart failure, Hypertension, Age 75+ years, Diabetes mellitus, Stroke, Vascular disease, Age 65-74 years,

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**Figure 4** A flow chart describing the clinical decision-making of the study cardiologist for participants following up in clinic for abnormal screening results during SMART-India study. CHA₂DS₂-VASc, Congestive heart failure, Hypertension, Age 75+ years, Diabetes mellitus, Stroke, Vascular disease, Age 65-74 years, Female sex; HbA1c, glycated haemoglobin; SMART-India, Smartphone Monitoring for Atrial fibrillation in Real-Time—India.
Female sex (CHA\textsubscript{DS\_2}-VASc) score.\textsuperscript{25} All of the clinical data including medications is abstracted by the clinical coordinator using clinical research forms.

**Modifications for follow-up scheduling**

A preliminary inspection of clinical follow-ups found that the majority are due to an ‘inconclusive’ finding of AliveCor ECG recording based on Kardia’s automated algorithm. Specifically, 44 out of the first 50 follow-ups are of participants whose screening was inconclusive. Out of these 44 participants, 41 were found to have normal sinus rhythm after the study cardiologist interpreted field and 12-lead clinical ECGs. Additionally, two of the remaining three participants were found to have benign, non-AF arrhythmias. Based on these findings, we modified our follow-up protocol by having the study cardiologist triage the single-lead ECG recordings from the field prior to the participants’ clinical follow-ups to prioritise high-risk patients and rule out false-positive recordings.

**Data adjudication**

All interpretations of ECG recordings determined as AF or abnormal by the AliveCor automated algorithm are reviewed by a separate team of trained clinicians through a standardised adjudication process. Two resident-level physicians independently review each abnormal AliveCor generated ECG recording using REDCap and enter their interpretation as either: (1) normal sinus rhythm, (2) AF or (3) as an other arrhythmia (free text). In the event of disagreement between the two reviewers, a fellow-level physician reviews the ECG recording.

**Analytical plan**

**Aim 1: AF epidemiology**

The gold standard of cardiologist-verified diagnoses of AF will be used to calculate the age-stratified and sex-stratified prevalence of AF in the study sample. Population estimates will be derived from the sample prevalence using survey weights constructed using the census collected by the SPARSH programme and the Government of India sponsored census in 2011. Three separate calculations will be performed to describe the overall prevalence of AF and separately for those with valvular AF and non-valvular AF.

**Aim 2: Profile of patients likely to have AF**

Distribution of covariates (table 1) across group of participants diagnosed with AF will be compared with those who were not diagnosed with AF to determine associated risk factors. Sensitivity analyses will be performed to rule out differences between the group of participants who were not diagnosed with AF but were followed up clinically in comparison with those who were never followed up. If meaningful differences are identified in our sensitivity analyses, distribution of covariates will be assessed across the three different groups of participants. Additionally, among those participants who are followed up clinically, differences in the distribution of clinical factors between the group of AF-positive and AF-negative participants will be assessed. For sociodemographic and clinical risk factors, multivariable regression techniques will be used to identify their conditional effects after accounting for other covariates. Lastly, among the participants who were followed up clinically, the effects of AF will be estimated by comparing differences in the quality of life score ((Short Form-12 Survey) SF-12, *table 1*) between groups of participants who are diagnosed for AF versus those who do not have AF. Potential confounders will be identified based on a priori knowledge (age, sex, socioeconomic status) and an examination of bivariate association of sociodemographic factors with AF and quality of life.

**Aim 3: Performance of AF-screening technology**

Sensitivity, specificity and discriminative ability (receiver operating characteristic curve analyses) with 95% CI bands will be used to measure the performance of our AF-screening technology in field and in clinic (table 4). In-field performance of AF-screening technology will be used by comparing results from the ANAND and AliveCor automated algorithms with the gold standard of adjudicated interpretations of the AliveCor ECG recordings. The clinical performance of AF-screening technology will be assessed by comparing results from the automated algorithms from both technologies with results from the gold-standard 12-lead ECG.

**Limitations and strengths**

It is important to consider that this study screens people of villages from rural Indian region who live in villages enrolled in SPARSH programme that is focused on addressing non-communicable diseases and its risk factors. Therefore, the participants recruited in this study may under-represent the true burden of AF risk factors present in rural Indian population. Additionally, ascertainment of certain lifestyle-related AF risk factors is based on self-report, which is prone to under-reporting. Mobile technologies are used to perform multiple AF

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**Table 4  Validation strategy to assess performance of atrial fibrillation screening technology**

<table>
<thead>
<tr>
<th>Phase</th>
<th>Screening technology</th>
<th>Gold standard</th>
</tr>
</thead>
<tbody>
<tr>
<td>In-field</td>
<td>Automated Novel Atrial fibrillation Noninvasive Detection (ANAND) automated algorithm AliveCor automated algorithm</td>
<td>Clinical adjudication of AliveCor recording from the same day</td>
</tr>
<tr>
<td>In-clinic</td>
<td>ANAND automated algorithm AliveCor automated algorithm</td>
<td>Clinical interpretation of 12-lead ECG from the same day</td>
</tr>
</tbody>
</table>
screening to account for potentially paroxysmal nature of AF. This approach, although superior to single time point screening, is vulnerable to missing AF. However, we attempt to partially address this limitation by obtaining a 24-hour Holter recording for participants who experience rhythm abnormalities on AliveCor or 12-lead ECG but may not be classified as AF.

DISCUSSION

The SMART-India study represents an innovative screening and referral paradigm for persons living in low-resource settings who may have an undetected chronic disease. Our study also highlights the importance of conducting a feasibility study and engaging local community partners prior to undertaking more broad-based population screening efforts. Findings from this study will provide greater insights into the silent burden of AF in rural Indian communities and help identify potential targets for intervention to modify the ongoing epidemic of stroke in India. The potential public health impact of the SMART-India study is particularly noteworthy in light of a recent white paper on AF screening which reported that AF detected during a screening procedure is not a benign condition and carries a sufficient risk of stroke. Further work is needed to build on the SMART-India study and build a nexus of providers that can help triage high-risk patients and further enhance the efficiency of a referral system for persons with suspected chronic disease, including cardiovascular disease. Leveraging existing community-based programmes, such as SPARSH, can help achieve this ambitious, but feasible, goal.

Ethics and dissemination

The SMART-India study was reviewed and approved by ethical boards from the University of Massachusetts Medical School, Worcester, Massachusetts, USA and the National Center for Advancing Translational Sciences (TL1-T1001454) and JAMA received support from the National Institute on Minority Health and Health Disparities (R63-MD008912-05). DDMM’s time was supported by KL2RR031981, 1R15HL121761-01A1, 1U54RR030952-01A2 and 1R01HL128911-01A1 from the National Heart, Lung and Blood Institute.

Disclaimer

The content is solely the responsibility of the authors and does not necessarily represent the official views of the NIH.

Competing interests

DDMM discloses equity stakes or consulting relationships with Flexcon, Inc., Bristol-Myers Squibb, Inc., Mobile Sense, Inc., ATRIA, Inc. and Boston Biomedical Associates, Inc. He has also received research funding from Sanofi Aventis, Inc., Otsuka Pharmaceuticals, Inc., Philips Healthcare, Inc., Biotronik, Inc. and Pfizer, Inc.

Patient consent

Obtained.

Ethics approval

UMMS IRB & CAM HREC.

Provenance and peer review

Not commissioned; externally peer reviewed.

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