Self-Screening and Non-Physician Screening for Hypertension in Communities: A Systematic Review

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BACKGROUND
Community-based self-screening may provide opportunities to increase detection of hypertension, and identify raised blood pressure (BP) in populations who do not access healthcare. This systematic review aimed to evaluate the effectiveness of non-physician screening and self-screening of BP in community settings.

METHODS
We searched the Cochrane Central Trials Register, Medline, Embase, CINAHL, and Science Citation Index & Conference Proceedings Citation Index—Science to November 2013 to identify studies reporting community-based self-screening or non-physician screening for hypertension in adults. Results were stratified by study site, screener, and the cut-off used to define high screening BP.

RESULTS
We included 73 studies, which described screening in 9 settings, with pharmacies (22%) and public areas/retail (15%) most commonly described. We found high levels of heterogeneity in all analyses, despite stratification. The highest proportions of eligible participants screened were achieved by mobile units (range 21%–88%) and pharmacies (range 40%–90%). Self-screeners had similar median rates of high BP detection (25%–35%) to participants in studies using other screeners. Few (16%) studies reported referral to primary care after screening. However, where participants were referred, a median of 44% (range 17%–100%) received a new hypertension diagnosis or antihypertensive medication.

CONCLUSIONS
Community-based non-physician or self-screening for raised BP can detect raised BP, which may lead to the identification of new cases of hypertension. However, current evidence is insufficient to recommend specific approaches or settings. Studies with good follow-up of patients to definitive diagnosis are needed.

Keywords: blood pressure; community health services; hypertension; primary care; screening; self-evaluation; self-screening.

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The aim of this study was therefore to systematically review the evidence for self-screening and other community-based non-physician screening strategies, and the associated rates of target populations screened, detection of raised BP, and rates of follow-up and intervention, including new diagnoses, in primary care. A further aim was to establish the factors, such as appropriate settings, screeners, and targeted populations that underpin a successful screening program for hypertension. We considered that a successful screening program would obtain good coverage of the targeted population, and refer all new cases of screen-detected high BP for definitive diagnosis without excessive numbers of false-positive referrals.

**METHODS**

**Search strategy**

We searched the Cochrane Central Register of Controlled Trials (Issue 4 2011), Medline (from 1948), Embase (from 1974), CINAHL (from 1980), and Science Citation Index & Conference Proceedings Citation Index—Science (from 1945) from the start of each database until 25th November 2013 using a search strategy developed with the assistance of an information specialist. The search strategy included terms relating to BP measurement, community sites and screeners, and self-screening (Supplementary Appendix A). We subsequently added an exclusion criterion for studies conducted prior to 1980, as initial screening confirmed that suitable devices for self-screening were not generally available before this date. Additional studies were identified by searching reference lists of included studies.

**Inclusion and exclusion criteria**

We included studies reporting self-screening or community-based screening by non-physicians of BP in adults ≥18 years of age (Table 1). “Self-screening” included home measurement of BP for screening purposes, but not for diagnostic or monitoring purposes. Screening carried out by clinically trained screeners (e.g., nurses) in clinical settings were excluded, although we included studies in community dentistry, as routine BP measurement is not usually practiced in community dentistry, and so this represents a novel setting for BP screening. No limit by comparator or study design was applied. Studies where BP measurements were obtained during screening with the primary aim of detecting a condition other than hypertension, as well as those where multiple screening tests were performed in a single visit were included.

**Study selection**

Two authors (S.F and H.A.) screened all titles and abstracts, and retrieved full text articles for potentially eligible studies. Where necessary, authors of studies were contacted up to 2 times by electronic mail for clarification of inclusion status. If study eligibility could not be confirmed after 2 attempts to contact the authors, the study was excluded.

**Data extraction**

Data from each paper were extracted by one author (S.F, H.A., D.M., R.J.M., C.H., or A.W.), and the data extraction form checked by a second author in the group, with disagreements resolved by consensus or a third author (S.F or H.A.). Extracted data included study characteristics, the number screened, and where reported, the proportion eligible for screening, the proportion with screen-detected hypertension, the hypertension threshold used, the proportion of existing hypertension, and any follow-up carried out.

**Assessment of quality**

Quality assessment was conducted using criteria based on the Quality Assessment tool for Diagnostic Accuracy Studies-2 (QUADAS-2) at the same time as extraction, and by the same authors, with disagreements resolved by consensus or a third author (S.F or H.A.). The criteria assessed applicability and bias due to patient selection, appropriateness of patients and setting, method of BP measurement, the prespecification and appropriateness hypertension threshold, and blind to preexisting hypertension status (Supplementary Appendix B).

**Data synthesis**

Characteristics of included studies were summarized, including the populations screened, setting, and aspects of study design (Supplementary Appendix C).
Data on the proportion of participants screened, and the proportion with high screening BP were reported grouped by the hypertension threshold defined in the study, size of study (number screened), qualification of person conducting the screening, and type of setting.

High levels of data heterogeneity meant that statistical synthesis was not possible. However, where at least 3 similar studies were included, their results were summarized using a box and whiskers plot, and possible factors relating to outliers discussed. We defined outliers as values further than 1.5 interquartile ranges from the nearest quartile. Statistical analyses were carried out using R (version 3.0.1, R Foundation for Statistical Computing).

RESULTS

We identified 16,356 studies (Figure 1). After initial screening of titles and abstracts, we examined 285 full text papers, of which 73 met our inclusion criteria.

Characteristics of included studies

An overview of the variation in study design and location is given in Table 2. Where reported (18 out of 73 studies, 25%), age of participants varied from a mean age of 41–75 years. Several studies targeted specific populations; for example, 14 studies screened ethnic minority populations, and 2 screened men.

Screening was conducted in 9 different types of setting (Table 2). The majority of studies reporting data on the number of screening sites used (n = 29 out of 49 studies reporting these data, 59%) described between 1 and 5 screening sites.

The most frequently reported types of people performing the screening were nurses (n = 17, 23%), followed by lay screeners (n = 16, 22%), and only 3 studies used self-screening (Table 2). Only 39 (53%) studies reported the type of BP measurement device used, which included automated devices in 25 (34%) studies, and a manual sphygmomanometer in 14 studies (19%). The number of measurements taken on each occasion was reported by 27 studies and ranged from 1 to 12. If more than one BP was taken there was significant variation in the timing of subsequent measurements.

Quality assessment

The results of the quality assessment (Figure 2) showed that few studies reported blinding to existing hypertension status (Supplementary Appendix D). Many studies (62%) were assessed as vulnerable to bias due to their recruitment.
methods, such as convenience sampling. The method for measuring BP was also poorly reported in the majority of studies, although most studies prespecified an appropriate threshold for hypertension.

**Proportion of eligible participants screened**

Twenty-three studies reported data for both the eligible population and the number screened, allowing the coverage of the screening strategies to be calculated. There was considerable heterogeneity both between and within sites (Figure 3): for example, 5 studies conducted in community buildings found that 12%–89% of eligible people were screened. A similar range was noted for screeners: for example, where nurses performed the screening, the proportion screened ranged from 5.5% to 99%. Mobile units (range 21%–88%) and pharmacies (range 40%–91%) achieved relatively high coverage across a range of studies. Some of the observed heterogeneity in coverage can be explained by differences in how eligible participants were defined in different studies. For example, 3 studies conducted in public areas and retail sites found that 5.5%–97% of eligible people were screened, the main differences being whether the population at large or the population...
Fleming et al.

attending a particular setting was defined as the denominator. Results also varied widely both within and between each type of screener. Student screeners achieved a low range of coverage (3.4%–36%), whereas pharmacy staff obtained higher coverage of the eligible population (56%–91%).
Detection of hypertension in screened participants

Fifty-nine studies contained sufficient data to calculate the proportion of screened participants who had hypertension. Although the definition of hypertension varied considerably between studies, there did not appear to be any relationship between hypertension threshold and detection rate (Supplementary Appendix F).

Most studies, whether grouped by site or screener, showed considerable heterogeneity in detection rate. Two types of site appeared to have lower detection rates: dental sites (range 8.5%–39%) and mobile sites (range 3.2%–34%) (Supplementary Appendix G). A number of the mobile sites served communities with reduced access to healthcare (e.g., immigrant communities in the United States).

It was notable that the 2 self-screening studies (i.e., those where participants measured their own BP)\textsuperscript{14,15} both resulted in detection rates (29% and 36%) comparable to those where participants measured their own BP to healthcare (e.g., immigrant communities in the United States).

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One outlier study\textsuperscript{16} was found during the analysis, with considerably lower detection rates than the other studies carried out in pharmacies. This paper was relatively old and used considerably higher hypertension cut-offs than more contemporary work (variable with minimum cutoff of 160/100 mm Hg), which would be expected to result in lower detection rates given a similar population.

Detection of hypertension in participants with and without preexisting diagnoses

Twenty-four studies provided data on detection of raised BP in participants with and without preexisting hypertension diagnoses. Figure 5 shows that in many of these studies, the proportion of participants with uncontrolled hypertension (i.e., those with an existing hypertension diagnosis and a raised BP on screening) equaled or exceeded the proportion of potential new diagnoses (i.e., participants with raised BP on screening, but without any known diagnosis of hypertension). Most of these studies recruited from populations which were likely to be representative of the general population in the study area, rather than selected subgroups.
Follow-up of participants after screening

Twelve studies provided data on participants following screening (Supplementary Appendix E). Only 3/7 (43%) referred all participants with elevated screening BP to primary care, and in one case, as few as 26% of those screening positive were referred. Eight studies reported the outcome of the referral, with a median of 44% (range 17%–100%) of those referred receiving a new hypertension diagnosis or new antihypertensive treatment. In 6 out of the 8 studies, this proportion exceeding 40%.

DISCUSSION

The results of this review show that community screening of BP by non-physicians (including self-screening) appears effective for detecting raised BP at screening, and that where data exists for this outcome, high screening BP leads to a new diagnosis or treatment for hypertension in a median of 44% of those referred. However, given the large amount of heterogeneity between study designs, it was not possible to carry out meta-analyses. Combined with the large amounts of variability in results between studies, this makes it difficult to recommend an optimal approach or setting. However, given the likely lower cost of self-screening compared to screening using nurses and other healthcare professionals, it is notable that self-screening appears to detect a similar proportion of participants with high BP as other screening strategies.

While many patients with existing hypertension may use community screening facilities, this is not necessarily a concern, particularly given the proportion of treated patients with uncontrolled BP. Raised pressure in treated hypertensives may prompt them to seek further appropriate medical care. Clear pathways for the referral and follow-up of patients identified with increased BP at screening are essential if the diagnosis of hypertension is to be established and any management benefits realized, but few studies reported any aspect of such follow-up. In many cases, raised screening BP was not subsequently acted upon, with only 3 out of 7 studies evaluating a system with universal referral of all such participants. Reassuringly, where participants were referred and subsequently attended primary care, around 40% received an intervention (new hypertension diagnosis or antihypertensive therapy), indicating the value of follow-up.

Considerable heterogeneity was observed across all analyses, despite stratification, precluding formal meta-analysis. The remaining heterogeneity likely reflects differences in the underlying study populations, and how well these populations are able to access hypertension diagnosis and treatment options. Additional potential sources of heterogeneity include methods of BP measurement, participant age range (which was typically poorly defined), and the influence of recruitment methods, some of which may have been more effective at targeting high-risk populations.

Blinding to existing hypertension status was poorly reported, but other than self-screening, there are no methodological barriers to blinding observers to this outcome. Although a lack of blinding would not be a significant concern for studies using automated devices, many studies in the review used manual BP measurement, which is known to be subject to terminal digit preference, and so observer bias may have been an issue for some of these studies where blinding was not present.17,18

The majority of studies (39, 53%) were undertaken in the United States. We may have missed studies from other countries; however, we did screen over 16,000 studies, and examined 285 full text papers. This suggests a lack of generalizability of our findings to other countries, as the effects of

Figure 5. Proportion of screened participants with and without preexisting hypertension diagnoses found to have high screening blood pressure.
screening may be dependent on the organization and coverage of the healthcare system.

No previous systematic reviews of community screening or self-screening for hypertension were identified in the search results, suggesting that this paper provides the first comprehensive review of the literature on this subject. International community-based surveys of BP and hypertension prevalence continue to suggest both significant proportions of undetected hypertension and of poor control among those detected.\textsuperscript{19}

The quality of the studies we found limits our ability to come to a definite conclusion as to the best method for implementing hypertension screening. We therefore perceive the need for high-quality studies that include BP target definitions, measurement methods, and subsequent data on the proportion of new hypertensive patients diagnosed. Indeed, those implementing self-screening should routinely embed a robust evaluation to underpin the ongoing use of resources in this area. To ultimately determine the cost effectiveness of these methods, there will be a need for robust randomized trials.

New developments in technology, including the use of social media to promote screening, use of smartphones, and enhanced network connectivity enabling screening results to be incorporated directly into electronic medical records, may improve the uptake and efficacy of community BP screening, and particularly self-screening.\textsuperscript{20}

As few studies followed up all participants to diagnosis, we are unable to assess what proportion of participants with raised BP at screening had true hypertension. The lack of studies with adequate follow-up make it currently impossible to determine whether these initiatives are effective or worthwhile. Properly powered studies including both cost and efficacy data are therefore required. As with standard clinical practice, such studies should follow-up all cases of raised BP with robust diagnostic methods.

In conclusion, despite a large number of studies reporting community-based BP screening by non-physicians, the evidence base for its effectiveness is very poor. Detection of high screening BP by self-screening is similar to that of more robust diagnostic methods, and efficacy data are therefore required. As with standard clinical practice, such studies should follow-up all cases of raised BP with robust diagnostic methods.

In conclusion, despite a large number of studies reporting community-based BP screening by non-physicians, the evidence base for its effectiveness is very poor. Detection of high screening BP by self-screening is similar to that of more robust diagnostic methods, and efficacy data are therefore required. As with standard clinical practice, such studies should follow-up all cases of raised BP with robust diagnostic methods.

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