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a total of 37 studies. Increasing the number of days of measurement improved prognostic power: 72%-91% of the theoretical maximum predictive value (asymptotic maximum hazard ratio) was reached by 3 days and 86%-96% by 7 days. Increasing beyond 3 days of

Schedules for Self-monitoring Blood Pressure: A Systematic Review

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BACKGROUND

Self-monitoring of blood pressure better predicts prognosis than clinic measurement, is popular with patients, and endorsed in hypertension guidelines. However, there is uncertainty over the optimal selfmonitoring schedule. We therefore aimed to determine the optimum schedule to predict future cardiovascular events and determine "true" underlying blood pressure.

METHODS

Six electronic databases were searched from November 2009 (updating a National Institute for Health and Care Excellence [NICE] systematic review) to April 2017. Studies that compared aspects of self-monitoring schedules to either prognosis or reliability/reproducibility in hypertensive adults were included. Data on study and population characteristics, self-monitoring regime, and outcomes were extracted by 2 reviewers independently.

RESULTS

From 5,164 unique articles identified, 25 met the inclusion criteria. Twelve studies were included from the original NICE review, making

Hypertension is a key risk factor for cardiovascular disease, the most important cause of morbidity and mortality worldwide.¹ The detection and subsequent management of hypertension requires appropriate monitoring, and selfmonitoring of blood pressure (SMBP) is increasingly used for

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this purpose with endorsement by guidelines worldwide.²⁻⁴ Compared to office blood pressure measurement, home readings better predict end organ damage, provide a more accurate diagnosis of hypertension, and improve patient involvement in their own care.5-7

measurement was necessary. CONCLUSIONS

Home blood pressure should be measured for 3 days, increased to 7 only when mean blood pressure is close to a diagnostic or treatment threshold. Other aspects of a monitoring schedule can be flexible to facilitate patient uptake of and adherence with self-monitoring.

measurement did not result in better correlation with ambulatory

monitoring. There was no convincing evidence that the timing or

number of readings per day had an effect, or that ignoring the first day's

Keywords: blood pressure; blood pressure monitoring; hypertension; regression dilution; schedule; self-monitoring; systematic review.

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Despite the growing popularity of SMBP, there is little agreement as to the optimal self-monitoring schedule. The Japanese Society of Hypertension guidelines recommend 2 readings on each occasion, using the mean of the 2 over 5–7 days.⁸ The European Society of Hypertension, along with the American Heart Association and the American Society of Hypertension and National Institute for Health and Care Excellence (NICE), recommend that blood pressure (BP) should be measured on at least 3–4 days and preferably on 7 consecutive days in the morning and evening, with 2 measurements per occasion taken 1–2 minutes apart. The readings taken on the first day should be discarded and then the average of the remaining readings used.^{2,3,9} There are no separate schedules recommended for ongoing management of patients with hypertension once the initial diagnosis has been made.

This study aimed to assess the evidence for these various guideline recommendations using the systematic search undertaken for the NICE (2011) Hypertension Guidelines² as a starting point, and updating and reappraising the literature.

METHODS

Data sources and searches

Electronic databases (Cochrane Central Register of Controlled Trials [The Cochrane Library, Wiley] (issue 3, March 2017), Medline [OvidSP] (1946–present, in process), Embase [OvidSP] (1974-present), CINAHL [EBSCOhost] (1980-present), Science Citation Index [Web of Knowledge] (1945-present), and Conference Proceedings Citation Index-Science [Web of Knowledge] (1945-present)) were searched up to April 2017, for articles published from November 2009 onward based on a search strategy developed for the NICE Hypertension Guidelines.² The original NICE search was of Medline, Embase, CINAHL, and the Cochrane Library from inception to November 2010 and the update search dates were chosen with some overlap to ensure relevant studies would not be missed. The search strategy for Medline can be found in Supplementary Appendix 1, which was then adapted for the other databases.

Study selection

Two reviewers independently reviewed the titles and abstracts of potentially relevant articles for inclusion. Full papers of potentially eligible articles resulting from the search plus all included articles from the NICE review were then assessed.

All study design types were eligible for inclusion. Studies must have assessed SMBP defined as BP measurement by a patient or carer, without the involvement of a health professional. It was anticipated that included studies would compare one or more of the following protocol components: number, timing, frequency, and duration of measurements and whether any readings should be discarded, but included all studies that compared any aspects of self-monitoring schedules. Studies that assessed regimes in terms of BP variability, machine validation studies, those containing inadequate description of the self-monitoring protocol, or Participants of interest were adults (18 years and older) with treated or untreated hypertension, who may or may not have had a comorbid disease. Reliability/reproducibility studies were included where at least some of the participants had hypertension or were being assessed to confirm suspicion of hypertension (e.g., where a previous clinic reading had indicated hypertension), and similarly prognostic studies (which were all conducted in the general population) where at least some participants either had hypertension or were treated with antihypertensive medication.

Articles written in a language other than English were translated to assess eligibility.

Data extraction and quality assessment

Data from each article were extracted independently by at least 2 reviewers using piloted forms (Supplementary Appendix 2). Information collected included study (e.g., country, hypothesis) and sample (e.g., sample size, age, comorbidities) characteristics, self-monitoring regime details (e.g., frequency, duration, whether devices used were validated), and outcome measures (see later). Any discrepancies were resolved by consensus.

A priori outcomes of interest varied with the type of study

- (1) Prognostic studies: mortality, stroke, myocardial infarction, angina, and heart failure or composites thereof.
- (2) Reliability/reproducibility studies: reproducibility of SMBP or correlations with ambulatory blood pressure measurement (ABPM) or office blood pressure measurement.

Methodological quality was assessed using an adaptation of 3 validated checklists: Effective Public Health Practice Project, Downs and Black, and Quality Assessment of Diagnostic Accuracy Studies (QUADAS-2).^{10–12} Additional questions about the validation status of the BP monitoring equipment used in each study were incorporated, for which we consulted the dabl Educational Trust and British and Irish Hypertension Society websites^{13,14} rather than rely on author-reported validation status (Supplementary Appendix 3 provides details of the methodological quality checklist applied).

Data synthesis and analysis

Imprecision in a measurement makes associations, such as hazard ratios (HRs) or correlations, harder to observe. Averaging over several measurements can reduce imprecision. Hence, a single imprecise measurement will show a weaker apparent association with an outcome, but increasing the number of measurements increases the apparent association.

To enable a consistent measure of comparison, the adjusted HR per 5 mm Hg increase in systolic BP was calculated for prognostic studies across the number of days of readings they considered. Study-specific curves for HR against number of days (n) were estimated by assuming that the reciprocal of estimated log HR was linear in 1/n, and for estimated correlation coefficient against *n* by assuming

that the reciprocal of correlation squared was linear in 1/n. These relationships were derived from standard results for linear regression dilution, which have been shown to apply approximately for HRs, under independence assumptions, when censoring is present and the sample size is large¹⁵ (Supplementary Appendix 4 provides a further explanation of the method of analysis used, including the approach to regression dilution). For each study, the "maximum log hazard ratio" was defined as the asymptotic maximum of the fitted curve on the log HR scale (i.e., the best log HR that could theoretically be achieved given an infinite number of days for measurement) and the fitted log HR at day 3 and day 7 is reported as a percentage of this maximum.

For reliability/reproducibility studies, the correlations reported between systolic and diastolic SMBP and ABPM as the reference standard were summarized. The remaining studies, in particular reliability studies that reported correlations with measures other than ABPM, were considered too dissimilar to group.

RESULTS

A total of 5,164 unique citations were identified of which 297 were assessed in detail along with 13 articles from the NICE search (Figure 1). Thirty-seven studies proved eligible for inclusion in the analysis comprising 25 from the update and 12 from the original NICE search (the remaining

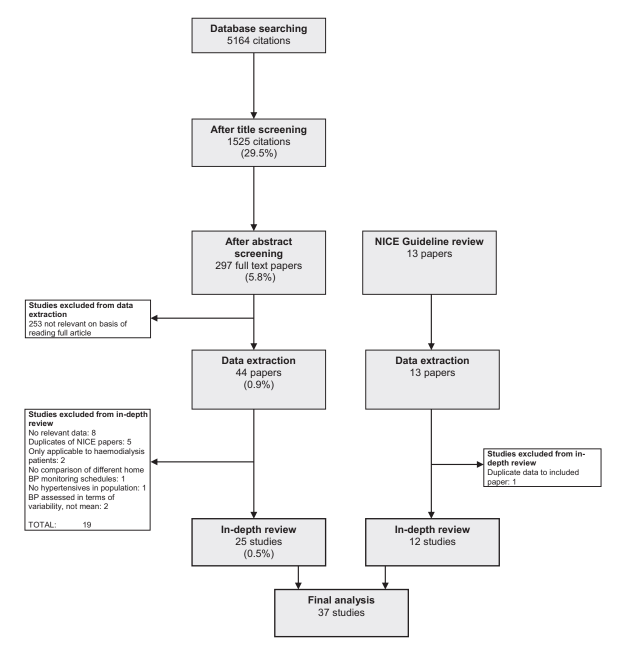


Figure 1. Filtering of papers from searching to synthesis.

Abbreviations: BP, blood pressure; NICE, National Institute for Health and Care Excellence.

article from the NICE review provided only duplicate data). Participants in the included studies (Supplementary Table 1) were drawn from 18 different countries and varied markedly in terms of mean age (range 40–70 years), gender (percentage male 26%–100%), sample size (43–21,591), and the proportion with hypertension and/or on antihypertensive medication (0%–100%).

Of the 37 articles, 10 were prognostic and 27 were reliability/reproducibility studies. The wide range of aspects of monitoring schedule assessed in the included studies is shown in Supplementary Table 2. Owing to the heterogeneity of the self-monitoring protocols, and the variability in the clinical outcomes and analyses in the eligible studies, meta-analysis was not possible.

Methodological issues

All studies had some degree of methodological flaw (or lack of clarity in what was reported), with 16 (43%) studies not clearly using validated devices throughout (Supplementary Table 1). Although selection criteria of participants were generally clear, only 16 (43%) studies used selection methods likely to avoid bias (Supplementary Table 2). Attrition reporting provided reasons for dropouts but typically not the characteristics thereof. Validation (from monitor memory or telemonitoring) of self-monitored readings was only clear and adequate in 10 studies. Reporting of results was generally adequate.

Prognostic studies

The 10 prognostic studies analyzed cohort data from Japan (Ohasama and home blood pressure measurement with Olmesartan Naive patients to Establish Standard Target blood pressure study [HONEST]), Finland (FINN Home), and Greece (Didima), or meta-analyzed data from Ohasama, FINN Home, and an additional Japanese cohort, Tsurugaya, which had not been published separately in a format we could extract relevant data from.^{16,17} There was overlap of populations within each cohort but differences in type of regime considered and/or the outcomes assessed. All participants in the prognostic studies were sampled from a general population. Three studies (Supplementary Table 1) had prediction of stroke/transient ischemic attack as the main outcome, whereas 4 used cardiovascular-related events, one considered both types of outcome separately, and 2 used composite cardiovascular end points including stroke.

Figure 2 shows the adjusted log HR per 5 mm Hg increase in systolic BP for each of the 5 studies (1 provided only unadjusted HRs with confidence intervals) that considered how outcome varied by length of monitoring in days (see also Table 1). HRs increased with additional days of readings across the studies with a flattening of the curves after 1 week for the 2 studies with longer follow-up and similar shaped curves for the shorter studies. However, confidence intervals overlapped between the most and least predictive measurement regimes (in terms of days of monitoring).

In Figure 2 the dotted line represents the maximum log HR for the 5 prognostic studies: 86%–96% of the maximum

Few data on the impact of time of day were available, but suggested that there was a maximum difference in HR of 0.09 per 5 mm Hg increase in systolic BP with overlapping confidence intervals between morning and/or evening measurements. There was also no convincing difference in prognostic ability when using the first and/or the second measurement on each occasion (Table 2).

Considering the total number of readings added little to the results for number of days, reflecting the limited data on readings per day (Supplementary Table 3). Only 1 prognostic study considered the effect of omitting first-day readings from the analysis, which made no difference to the HR (Table 3).

Analysis of reliability/reproducibility studies

Participants in the 27 reliability/reproducibility studies were largely either treated or untreated patients with hypertension, though populations ranged from heart transplant recipients and renal outpatients to company volunteers and attendees at a health education program (Supplementary Table 1). Three studies shared populations with the Japanese and Finnish prognostic studies.

Of the 20 studies considering reliability/reproducibility, 15 reported correlations with ABPM as the reference standard, 8 using mean daytime ABPM, 5 using 24-hour ABPM, and 2 using both daytime and 24-hour ABPM. These 15 studies were included in the remainder of the analyses.

The correlation between cumulative mean home systolic BP and diastolic blood pressure from 1 to 7 days of monitoring with ABPM as the comparator measurement is shown in Figure 3 (analysis restricted to those studies (n = 5) with correlations for at least 3 different counts of days; the dotted line represents the maximum correlation coefficient) and Table 4. Here the curves were very flat and there was no convincing increase in correlation after the fourth day of monitoring. Better than 90% of the maximum correlation with ABPM was achieved by 3 days. In many correlation studies, numbers of participants were small and confidence intervals were wide, but this pattern was observed even in larger studies (n = 464).

Data could only be extracted from 3 studies to assess the relationship between correlation with ABPM and the number of readings on each occasion, time of day of 2 readings (Figure 4 and Table 5), and total number of measurements (Table 6). As for the prognostic studies, varying the number of readings on each occasion and time of day of readings appeared to have little impact, while examining the effect of number of measurements overall again largely replicated the results for number of days. Similarly, discarding the readings from the first day of home monitoring made little difference to correlation with ABPM, whether readings more than 3 days or 1 week were being considered (Table 3).

Three studies considered particular aspects of monitoring schedules uniquely—the time interval between readings, a schedule including before-morning micturition and afternoon readings vs. 1 involving post-morning micturition

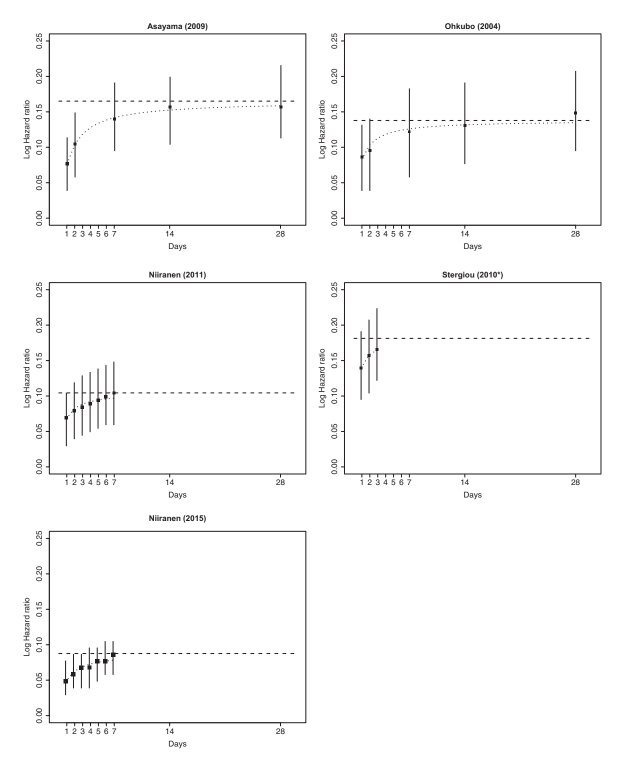


Figure 2. Log hazard ratios per 5 mm Hg for prognostic outcomes by cumulative numbers of days of self-monitoring. *Values from Stergiou (2010) are unadjusted hazard ratios (HRs).

and evening readings, and resting for 5 minutes before readings vs. not resting. However, the complexity of the schedule comparison in 1 study and the small sample size of the other studies prevented drawing any firm conclusions.

No study provided evidence on the timings of readings in relation to medications, how frequently monitoring should be repeated, or on whether fewer readings may be required for routine ongoing management.

author; publication Years of date) follow-up N(e Asayama (11.9, N(e (2009) ^a (median) 2 Ohasama 8.3, 4 Nirranen 8.3, 4 Nirranen Not stated 5 Muttiple Not stated 2 Not stated 2 Not stated 4						Adjustee	d HR per 5 mm	Adjusted HR per 5 mm Hg increase in systolic BP (95% Cl)	systolic BP (9	5% CI)		
a (median) (median) 8.3 (median) Not stated Not stated	R N (events) p	Readings per day	Outcome	1 day	2 days	3 days	4 days	5 days	6 days	7 days	14 days	4 weeks
8.3 (median) Not stated Not stated	2,234 (226)	1 daily (1 in the evening)	Stroke and TIA	1.08 (1.04–1.12)	1.11 (1.06–1.16)					1.15 (1.10–1.21)	1.17 (1.11–1.22)	1.17 (1.12–1.24)
b Not stated b Not stated Not stated	4,802 (568)	1 daily (1 in the morning)	CVD events	1.05 (1.03–1.08)	1.05 (1.03–1.08) (1.04–1.09)	1.07 (1.04–1.09)	1.07 (1.04–1.10)	1.08 (1.05–1.10)	1.08 (1.05–1.10) (1.06–1.11)	1.09 (1.06–1.11)		
	5,030 (588)	1 daily (1 in the morning)	CVD events							1.09 (1.06–1.11)		
	2,762 (360)	1 daily (1 in the morning)	CVD events								1.11 (1.07–1.15)	
	4,225 (509)	2 daily (1 in the morning and evening)	CVD events							1.11 (1.08–1.14)		
Niiranen (2011) 6.8 2 FINN Home (median)	2,081 (162)	4 daily (2 in the morning and evening)	CVD events	1.07 (1.03–1.11)	1.07 (1.03–1.11) (1.04–1.13)	1.09 (1.05–1.14)		1.09 1.10 1.10 1.10 1.10 (1.05–1.14) (1.06–1.15) (1.06–1.16)	1.10 (1.06–1.15)	1.11 (1.06–1.16)		
Ohkubo 10.6 (mean) 1 (2004) ^a Ohasama	1,491 (136)	1 daily 5 (1 in the morning)	Stroke and TIA	1.09 1.10 (1.04–1.14) (1.04–1.15)	1.10 (1.04–1.15)					1.13 (1.06–1.20)	1.14 1.16 (1.08–1.21) (1.10–1.23)	1.16 (1.10–1.23)
Stergiou 8.2 (mean) 662 (67) (2010) ^c Didima		4 daily (2 in the morning and evening)	CVD events	1.15		1.18 (1.13–1.25)						
Abbreviations: BP, blood pressure; CI, confidence interval; CVD, cardiovascular disease; TIA, transient ischemic attack. The references in the tables are cited in Supplementary Table 1. ^a All 3 studies from the same population, over slightly different time periods with slightly different focus—morning only, evening only, and morning and evening. ^b These 2 studies both use 3 datasets (including Ohasama and FINN Home). The 2013 study is an abstract before the main paper in 2015, but includes some different analyses. ^c Study reported upadilisted HRs without CIs (inst <i>P</i> < 0.05) Values in italics are unadilisted HRs. CIs taken from secondary paper in 2016, but includes also available from secondary paper.	essure; Cl s are cited e populatic 3 datasets HRs witho	l, confiden i in Supple on, over sl (including	ce interval; ementary Ta ightly differe J Ohasama st P < 0.05)	CVD, cardiov ble 1. ant time perio and FINN Ho	/ascular disea ds with slightl me). The 201 lics are unadi	CVD, cardiovascular disease; TIA, transient ischemic attack. ble 1. ent time periods with slightly different focus—morning only, ev and FINN Home). The 2013 study is an abstract before the m Values in italics are unadiusted HRs. Cls taken from second	ant ischemic a s-morning o bstract before	attack. nly, evening o the main pap	nly, and morn er in 2015, bu ar ¹⁸ Adiusted	CVD, cardiovascular disease; TIA, transient ischemic attack. ble 1. ent time periods with slightly different focus—morning only, evening only, and morning and evening. and FINN Home). The 2013 study is an abstract before the main paper in 2015, but includes some different analyses. Values in italics are unadiusted HRs. CIs taken from secondary paper in 2016, but includes also available from secondary	g. e different ana able from serv	Ilyses. andarv paper

6 American Journal of Hypertension

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	מ וומדמו מ ומנוי									
				1		Adjust	ed HR per 5 mm Hg	Adjusted HR per 5 mm Hg increase in systolic BP (95% Cl)	c BP (95% CI)	
Study.					Number of	Number of readings on each occasion	occasion		Time of day	
(first author; publication date)	Years of follow-up	N (events)	Readings schedule	Outcome	All first measurements	All second measurements	All measurements	All morning measurements	All evening measurements	All measurements Other
Asayama (2006) Ohasama	10.6 (median)	1,766 (156)	2 daily (1 AM and 1 PM) for 4 weeks	Stroke and TIA				1.14 (1.09–1.21)	1.14 (1.09–1.21) 1.16 (1.10–1.22) 1.17 (1.10–1.23)	1.17 (1.10–1.23)
Hoshide (2016) Ohasama	4.0 (mean)	4,278 (74) 4,278 (77)	4.0 (mean) 4,278 (74) 6 daily (3 AM and 3 PM) 4,278 (77) for 14 days	Stroke CVD events				1.17 (1.09–1.25) 0.96 (0.89–1.05)	1.17 (1.09–1.25) 1.12 (1.04–1.22) 1.18 (1.09–1.28) 0.96 (0.89–1.05) 1.05 (0.96–1.14) 1.00 (0.92–1.10)	1.18 (1.09–1.28) 1.00 (0.92–1.10)
Niiranen (2013) ^a Multiple studies	Not stated	5,030 (588)	1 daily (1 AM) for 7 days	CVD events				1.09 (1.06–1.11)		
	Not stated	4,225 (509)	2 daily (1 AM and 1 PM) for 7 days	CVD events						1.11 (1.08–1.14)
Niiranen (2011) FINN Home	6.8 (median)	2,081 (162)	4 daily (2 AM and 2 PM) for 7 days	CVD events	1.10 (1.06–1.15)	1.11 (1.07–1.16)	1.11 (1.06–1.16)	1.10 (1.06–1.15)	CVD events 1.10 (1.06–1.15) 1.11 (1.07–1.16) 1.11 (1.06–1.16) 1.10 (1.06–1.15) 1.10 (1.06–1.15) 1.11 (1.06–1.16)	1.11 (1.06–1.16)
Saito (2016) HONEST	2.0 (mean)	21,591 (280)	4 daily (2 AM and 2 PM) for 2 days over 7 periods	CVD events	CVD events 1.21 (1.15–1.27) 1.20 (1.14–1.27) 1.21 (1.15–1.27)	1.20 (1.14–1.27)	1.21 (1.15–1.27)			
Stergiou (2010) ^b 8.2 (mean) Didima	8.2 (mean)	662 (67)	662 (67) 4 daily (2 AM and 2 PM) for 3 days	CVD events	1.18	1.19	1.18 (1.13–1.25)	1.18	1.17	1.18 (1.13–1.25)
Abbreviations: BP, blood pressure; CI, confidence interval. Standard Target blood pressure; TIA, transient ischemic attacl Kario (2016) reports HRs for blood pressure categories for ^a This study uses 3 datasets (including Ohasama and FINN ^b Study reported unadjusted HRs without CIs (just $P < 0.0$. Cardiovascular risk prediction based on home blood pressure for all measurements only.	BP, blood pre blood pressure ports HRs for as 3 datasets (a unadjusted sk prediction b ints only.	essure; CI, s; TIA, tran: blood pres (including (HRs witho ased on hr	Abbreviations: BP, blood pressure; CI, confidence interval andard Target blood pressure; TIA, transient ischemic attacl Kario (2016) reports HRs for blood pressure categories for ^a This study uses 3 datasets (including Ohasama and FINN ^b Study reported unadjusted HRs without CIs (just $P < 0.0$ ardiovascular risk prediction based on home blood pressure ^a all measurements only.	val; CVD, cardi tack. for time of day, I NN Home). The 0.05).Values in ure measureme	ovascular diseast rather than per 1/. paper is an abstr. italics are unadju nt: the Didima str.	e; HONEST, hom 5/10 mm Hg, and act before the ful usted HRs. Cls t idy. J Hypertens:	le blood pressure I so cannot be incl I paper in 2015, bu aken from secon 2007; 25: 1590–1	Abbreviations: BP, blood pressure; CI, confidence interval; CVD, cardiovascular disease; HONEST, home blood pressure measurement with Olmandard Target blood pressure; TIA, transient ischemic attack. Kario (2016) reports HRs for blood pressure categories for time of day, rather than per 1/5/10 mm Hg, and so cannot be included in the above table. *This study uses 3 datasets (including Ohasama and FINN Home). The paper is an abstract before the full paper in 2015, but data in this table were ^b Study reported unadjusted HRs without CIs (just <i>P</i> < 0.05).Values in italics are unadjusted HRs. CIs taken from secondary paper: Stergiou GS reliovascular risk prediction based on home blood pressure measurement: the Didima study. <i>J Hypertens</i> 2007; 25: 1590–1596. Adjusted HRs also as all measurements only.	; CVD, cardiovascular disease; HONEST, home blood pressure measurement with Olmesartan Naive patients to E k. k. time of day, rather than per 1/5/10 mm Hg, and so cannot be included in the above table. Home). The paper is an abstract before the full paper in 2015, but data in this table were only published in this form. b5).Values in italics are unadjusted HRs. CIs taken from secondary paper: Stergiou GS, Baibas NM, Kalogeropou measurement: the Didima study. <i>J Hypertens</i> 2007; 25: 1590–1596. Adjusted HRs also available from secondary p	Abbreviations: BP, blood pressure; CI, confidence interval; CVD, cardiovascular disease; HONEST, home blood pressure measurement with Olmesartan Naive patients to Establish andard Target blood pressure, TIA, transient ischemic attack. Kario (2016) reports HRs for blood pressure categories for time of day, rather than per 1/5/10 mm Hg, and so cannot be included in the above table. [•] This study uses 3 datasets (including Ohasama and FINN Home). The paper is an abstract before the full paper in 2015, but data in this table were only published in this form. [•] Study reported unadjusted HRs without CIs (just <i>P</i> < 0.05). Values in falics are unadjusted HRs. CIs taken from secondary paper: Stergiou GS, Balbas NM, Kalogeropoulos PG. reliovascular risk prediction based on home blood pressure measurement: the Didima study. <i>J Hypertens</i> 2007; 25: 1590–1596. Adjusted HRs also available from secondary paper but reliovascular paper on home blood pressure measurement: the Didima study. <i>J Hypertens</i> 2007; 25: 1590–1596. Adjusted HRs also available from secondary paper on the secondary paper of the firm secondary paper of the above table.

Chudu (first suthan sublication data)	Commentar	N		Omitting measurements
Study (first author; publication date)	Comparator	N	Using all measurements	from the first day
3 days of home measurement			Correlation with ABPM (95%	confidence interval)
Johansson (2010)	24-hour ABPM	464	0.88 (0.86–0.90)	0.89 (0.87–0.91)
Stergiou (1998)	Daytime ABPM	189	0.68 (0.60-0.75)	0.67 (0.58– 0.74)
Verberk (2006)	Daytime ABPM	216	0.60 (0.51–0.68)	0.60 (0.51–0.68)
	24-hour ABPM	216	0.66 (0.58–0.73)	0.69 (0.61–0.75)
4 days of home measurement			Correlation with ABPM (95%	% confidence interval)
Di Monaco (2016)	Daytime ABPM	310	0.59 (0.51–0.65)	0.57 (0.49–0.64)
	24-hour ABPM	310	0.59 (0.51–0.66)	0.57 (0.49–0.64)
Stergiou (1998)	Daytime ABPM	189	0.70 (0.62-0.77)	0.69 (0.61–0.76)
Verberk (2006)	Daytime ABPM	216	0.62 (0.53-0.70)	0.62 (0.53-0.70)
	24-hour ABPM	216	0.68 (0.60-0.75)	0.69 (0.61–0.75)
1 week of home measurement ^a			Correlation with ABPM (95%	% confidence interval)
Johansson (2010)	24-hour ABPM	464	0.89 (0.87-0.91)	0.87 (0.85– 0.89)
Nunan (2015)	Daytime ABPM	203	0.67 (0.59–0.74)	0.68 (0.60-0.75)
Stergiou (1998)	Daytime ABPM	189	0.71 (0.63–0.77)	0.71 (0.63–0.77)
Verberk (2006)	Daytime ABPM	216	0.65 (0.57–0.72)	0.65 (0.57-0.72)
	24-hour ABPM	216	0.70 (0.62–0.76)	0.71 (0.64–0.77)
1 week of home measurement ^a			Hazard ratio for future CVD (9	5% confidence interval)
Niiranen (2011)	Future CVD	162 ^b	1.11 (1.06–1.16)	1.11 (1.06–1.16)

Abbreviations: ABPM, ambulatory blood pressure measurement; CVD, cardiovascular disease.

^a 1 week refers to 7 days of home measurement, except for Stergiou (1998) where home monitoring was only conducted for 6 days. ^b162 CVD events.

DISCUSSION

Summary of main findings

The literature has been comprehensively reviewed, finding 37 studies relating self-monitoring regimes to prognosis and/ or correlation to reference standard, with the aim of making evidence-based recommendations for future practice. For prognostic studies, only a small increase in precision was gained from undertaking more than 3 days of readings and the results from correlation studies were similar. Such differences are likely only to impact on clinical decision making around diagnostic or treatment thresholds. There was no convincing difference in terms of how many readings were taken per day, whether morning and/or evening measures are used, or whether the first day was removed.

Strengths and limitations of the study

This review used a comprehensive search strategy in multiple databases and all languages, incorporating hand searching, and is unlikely to have missed relevant articles. A thorough assessment of methodological quality was undertaken including assessment of the validation status of the monitors used.^{13,14} By estimating study-specific curves for either HR or

correlation coefficient against regime, the available data were synthesized in a robust form, despite any heterogeneity. By including a broad range of potential elements of monitoring schedules, this provides the most complete evidence to date on which to base recommendations.

The key weakness of this review is the paucity of studies of prognosis. Despite several different publications, only 4 sources of participants make up the full data set. While covering populations from Japan, Finland, and Greece, these data are lacking large relevant populations, in particular of South Asian and African/African Caribbean origin. Though used in a combined population, 1 cohort (Tsurugaya)^{16,17} was included, which has not been published separately in a format we could extract the relevant data from, and hence was only included as part of the Niiranen et al. meta-analysis.

Furthermore, the findings of small differences in both prognostic ability and correlation between different regimes must be tempered by the heterogeneity of design and methodological flaws identified in some studies. This reflected a lack of uniformity of method used between studies, and precluded comparison of more diverse regimes of measurement across multiple studies. Similarly, several studies used unvalidated equipment (Supplementary Table 1).

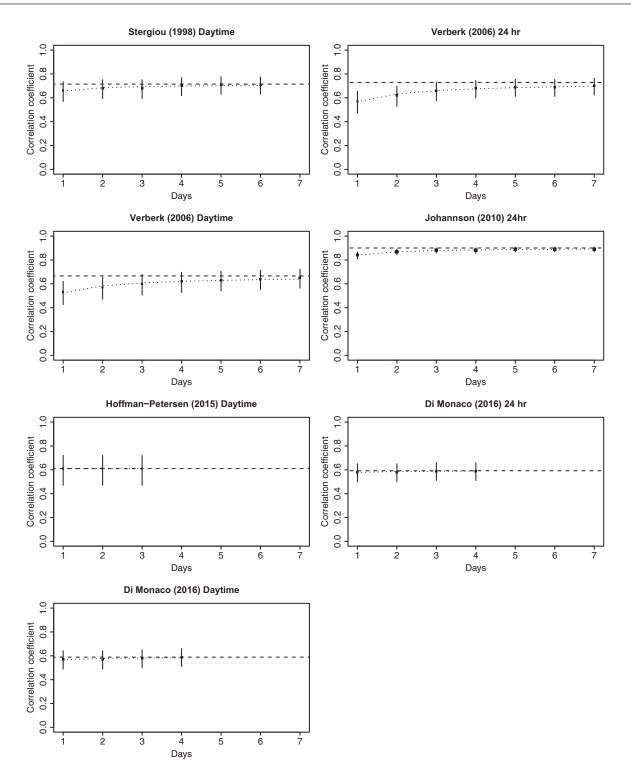


Figure 3. Correlation coefficient between ambulatory blood pressure measurement (ABPM) and self-monitoring of blood pressure (SMBP) by cumulative number of days of self-monitoring.

Comparisons with existing literature

One previous systematic review over a decade ago including 4 reliability/reproducibility studies¹⁹ considered multiple aspects of monitoring schedules but did not include any prognostic studies. In comparison, the current analysis includes 10 prognostic studies and 27 reliability/reproducibility studies. More recently, Niiranen et al.²⁰ combined 3 cohorts (2 Japanese and 1 Finnish) with consideration of prognosis in terms of number of days per week, but did not assess correlation data or other aspects of a monitoring regime, as the current work has.

author; publication Measurement date) schedule Almeida 3 days Before-morning (2014) micturition and in afternoon: 15 (3 day 1, 6 on days 2 and 3) adays post-morning micturition and evening: 15 (3 day 1, 6 on days 2 and 3) adays and 37 readings (2013) (9 on day 1, 12 on each of day 2 and day 3) Ambrosi 7 days (2 readings in con days 2 and (2014) 3 readings in the evening on each day) Boivin 3 readings without rest, 1.e., immediately after positioning cuff (Am and PM for 3 days, total 18 readings) 3 readings 5 minutes and PM for 3 days, total 18 readings in the conting and 2 readings in the evening on each day) Boivin 3 readings in the evening on each day) Boivin 3 readings in the evening on each day) Boivin 4 days (2 readings) Teadings in the conting and 2 readings in the evening on each day) Boivin 4 days (2 readings in the 2 re				Correl	Correlation coefficient between home systolic/diastolic BP and the comparator measurement	between hom	e systolic/dias	tolic BP and th	e comparator n	neasurement	
date) Schedule Almeida 3 days Before-mornir afternoom: 15 (3 day 6 on days 2 and 3) adys post-morning micturition and evening: 15 (3 day 6 on days 2 and 3) micturition and evening: 15 (3 day 6 on day 1, 12 on day 3) Almeida 3 days and 33 readin (3 on day 1, 12 on day 3) Ambrosi 7 days and 27 readin (3 on day 1, 16 on d day 3) Ambrosi 7 days (2 readings in the morning and 2 2-5) Ambrosi 7 days (2 readings in the morning and 2 2-5) Boivin 3 readings without rei (2014) Robin 3 readings in the evening on the and positioning cuff (AM and PM for 3 days, total 18 readings in the 2 readings in the evening on each d 2 readings in the			1	Time between readings	ß	Number o	of days (from f	Number of days (from first day onwards unless otherwise stated)	ds unless other	wise stated)	
3 3 3 3 3 3 3 4 4 4 4 4 4 4 4 4 4 4 4 4		Comparator	2	10 seconds 1 minute	ite 1 day	2 days	3 days	4 days	5 days	6 days	7 days
3 3 3 3 4 4 6 9 4 6 9 9 9 9 9 9 9 9 9 9 9 9 9 9	-morning ind in 5 (3 day 1, and 3)	24-hour wakefulness ABPM	158				0.769/0.826				
3 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	orning Ind (3 day 1, and 3)	Before SMBP					0.722/0.742				
5 (3 1 3 1 3 1 4 (4 (readings 12 on 2 and	24-hour ABPM	158				0.76/0.80				
7 (31 31 4 (readings 6 on days	Day before SMBP							0.61/0.69		
31 31 4 (ings in and 2 the each day)	24-hour ABPM Within 15 days end of SMBP	58				0.75/0.62			0.71/0.65 (Days 2–7)	
31	nout rest, ately after cuff (AM days, dings)	Daytime ABPM	52				99.0/69.0				
040	ninutes ries (∧M days, dings)	Within 3 days before SMBP					0.70/0.66				
4 0	ings in and	Daytime ABPM Day before SMBP	310		0.57/0.72	0.57/0.72	0.58/0.72	0.59/0.72			
4	n the each day)	24-hour ABPM Day before SMBP			0.58/0.71	0.58/0.71	0.59/0.71	0.59/0.72			
	ek for 8 adinas in th	Daytime ABPM	56	0.712/0.693 0.725/0.673	673						
morning and 3 in the evening on each day)	adungs in un 13 ng on	ABPM on second visit during SMBP schedule									
Hoffman- 4 days. 3 _{AM} (6–8 _{AM}), Petersen before dinner (2015) (5–7 _{PM}), 3 before bedtime (9–11 _{PM})	3–8 AM), 3 ∋r ⊃efore 11 PM)	Daytime ABPM Just after completion after SMBP	102		0.61/0.56	0.61/0.55 (days 2–3)	0.69/0.61 (days 2–4)				
Johannson 7 days (2 readings (2010)	ings ng and	24-hour ABPM	464		0.84/0.82	0.87/0.84	0.88/0.85	0.88/0.86	0.89/0.86	0.89/0.87	0.89/0.87
2 readings in the evening on each day)	n the each day)	Timing in relationship to SMBP unclear				0.89/ 0.87 (days 2–3)				0.87/ 0.85 (days 2–7)	

Important of the properties of the proproperties of the properties of the properties of the properties	Study (first	st			Corre	Correlation coefficient between home systolic/diastolic BP and the comparator measurement	between hom	e systolic/dias	tolic BP and the	e comparator n	neasurement	
Image: constant constan	author; publicatio	on Measurement		I	Time between readin	ßs	Number	of days (from fi	irst day onward	Is unless other	vise stated)	
Target Greening in the Strength in the	date)	schedule	Comparator	z			2 days	3 days	4 days	5 days	6 days	7 days
T days <i>L</i> readings Daytime ABM BT T days <i>L</i> readings ABM and SkiBP R manages in the second method of a solution table device or aBM first and some accord accord aBM first and some accord accord aBM first and some accord abm first and so	Kim (2015) [£]		24-hour ABPM	266				0.80		6.70		0.79
6495 (3 wheekens) Dayma ABPM 240 6496 (3 mod (0) w) Superingined day Superingined day 840 (3 mod (0) w) Superingined day Superingined day 1 are complexing Dayma ABPM 203 1 are complexing within game Dayma ABPM first, and (0) with a mod (0) with	McGowal (2010)	~	Daytime ABPM ABPM and SMBP within total 8-day period (i e., ABPM 1 day before or after SMBP)— some ABPM first and some SMBP first	87						0	0.72/0.89 (days 2–7)	
T days (t readings in the momenta) Daytime ABPM 203 Constrained (a)	Muxfeldt (2015)	5 days (3 AM between 6 and 10 AM) and 3 PM between 6 and 10 PM)	Daytime ABPM SMBP initiated day after ABPM	240					0.68/0.73 (days 2–5)			
3 work days per week for 2 weeks. Dayline ABPM 189 0.680.77 0.680.75 0.700.77 0.710.78 for 2 weeks. creatings in the morning and comming and some SMBP first. 0.880.77 0.680.77 0.690.77 0.710.79 0.710.79 7 days 13 readings in the worning on the worning of the wor	Nunan (2015)	7 days (2 readings in the morning and 2 in the evening)		203					0.658/0.707 (days 2–5)	0.649/0.703	0.68/0.71 (days 2–6)	0.671/0.708
7 days (3 readings in the morning and serial most in the morning and serial most in the averal most in the averal most in the averal most in the first of each inplicate discarded) 216 0.53/0.56 0.63/0.65 0.63/0.66 0.65/0.65 0.65/0.66 0.65/0.66 0.65/0.66 0.66/0.67 0.66/0.66 0.66/0.67	Stergiou (1998)	3 work days per week for 2 weeks (2 readings in the morning an the 2 readings in the evening on each day)	Daytime ABPM Some ABPM first and some SMBP first, one after the other	189		0.66/0.70	0.68/0.73 0.67/0.76 (days 2-3)		0.70/0.77 0.71/0.79 (days 2–5)	0.71/0.78 0.71/0.79 (days 2–6)	0.71/0.79	
216 0.57/0.61 0.62/0.63 0.66/0.66 0.68/0.66 0.69/0.69 (0.66/0.64 0.69/0.66 0.69/0.69 0.69/0.69 (1 days 2-4) (1 days 2-5) (1 days 2-5) (1 days 2-6) (1 days 2-6) (1 days 2-6) (1 days 2-6) (1 days 3-5) (1 days 3-7) (1 days 3-5) (1 days 3-6) (1 days 3-7) (1 days 3-5) (1 days 3-5) (1 days 3-7) (1 days 3-5) (1 days 3-7) (1 days 3-5) (1 days 3-7) (1 days 3-7) (1 days 3-5) (1 days 3-7) (1 days 3-5) (1 days 3-5) (1 days 3-7) (1 days 3-5) (1 days 3-5) (1 days 3-7) (1 days 3-5) (1 days 3-5) (1 days 3-7) (1 days 3-5) (1 days 3-7) (1 days 3-5) (1 days 3-5) (1 days 3-7) (1 days 3-5) (1 days 3-5) (1 days 3-7) (1 days 3-7) (1 days 3-5) (1 days 3-7) (1 days 3-5)	Verberk (2006)	7 days (3 readings in the morning and 3 readings in the evening, with the first of each triplicate discarded)	Daytime ABPM Timing vs. SMBP unclear	216		0.53/0.59	0.57/0.61 0.60/0.61 (days 2–3)		0.62/0.63 0.64/0.66 (days 2–5) 0.66(0.67 (days 3–6) 0.66(0.67 (days 4–7) (days 4–7)	0.63/0.66 0.65/0.66 (days 2–6) 0.66/0.67 (days 3–7)	0.65/0.66 0.65/0.66 (days 2–7)	0.65/0.66
			24-hour ABPM Timing vs. SMBP unclear	216		0.57/0.61	0.62/0.63 0.66/0.64 (days 2–3)	0.66/0.66 0.69/0.66 (days 2.41) 0.71/0.69 (days 3.5) 0.70/0.69 0.70/0.69 0.69/0.69 0.69/0.69 0.69/0.69 0.69/0.69 0.69/0.69 0.69/0.69 0.69/0.69	0.68/0.66 0.69/0.69 (days 2–5) 0.71/0.29 (days 3–6) 0.71/0.70 (days 4–7)	0.69/0.69 0.69/0.69 (days 2–6) 0.71/0.69 (days 3–7)	0.69/0.69 0.70/0.69 (days 2-7)	0.70/0.69

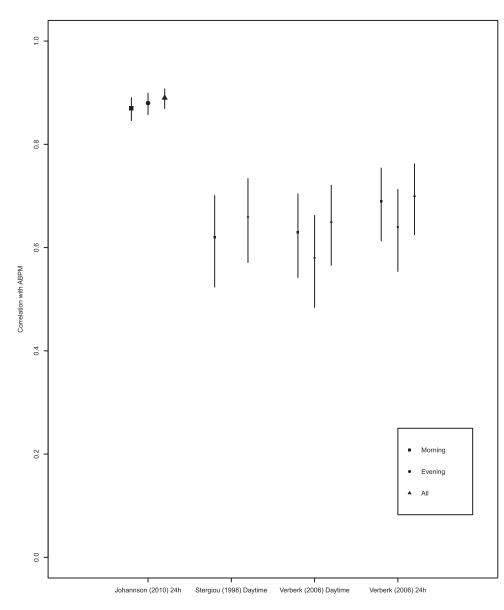


Figure 4. Correlation coefficient between ambulatory blood pressure measurement (ABPM) and self-monitoring of blood pressure (SMBP) by time of day of self-monitoring. The comparison in Stergiou (1998) is between first morning vs. first day; all other comparisons are between all morning, all evening or all readings.

Despite the heterogeneity and variable methodological quality of the evidence reviewed, many authors of the individual included studies drew strong and clear conclusions from their results. This was in spite of many HRs fully overlapping between apparently optimum and less optimum regimes. Subsequent guidelines from NICE, Europe, and the United States followed these conclusions in terms of recommendations on the number of days, the number of readings to take on each occasion, a preference for measuring in both the morning and evening, which values to discard, and total number of measurements.^{2–4} However, we found that for most aspects of monitoring schedules, evidence was either missing or at best ambivalent, suggesting excessive influence of the interpretation of individual studies by the study authors rather than the observed results.

Linked qualitative work by our group suggests that patients value flexibility in regime, and given the lack of evidence underpinning fixed regimes, incorporating such flexibility in future guideline iterations seems sensible.²¹ This might increase uptake and compliance,²² thus facilitating further implementation of self-monitoring.

Implications for clinical practice

The relatively modest benefit from more than 3 days of readings or of any particular quantity or timing of readings within these 3 days suggests that more protracted schedules are only likely to be worthwhile around diagnostic or treatment thresholds. Given the widespread use of telemonitoring, automated patient feedback could be used to

					Correlation co	sefficient betwee	n home systolic	Correlation coefficient between home systolic/diastolic BP and the comparator measurement	the comparator i	measurement
Study (first author:					Number of r	Number of readings on each occasion	occasion		Time of day	
publication					All first	All second	AII	All morning	All evening	AII
date)	Measurement schedule	dule	Comparator	z	measurements	measurements	measurements	s measurements	measurements	measurements
Johannson (2010)	7 days (2 readings in the morning and 2 readings in the evening on each day)	ay)	24-hour ABPM Timing in relationship to SMBP unclear	464	0.89/0.87	0.89/0.87	0.89/0.87	0.87/0.85	0.88/0.87	0.89/0.87
Stergiou (1998)ª	3 work days per week for 2 weeks (2 readings in the morning and 2 readings in the evening on each day)		Daytime ABPM Some ABPM first and some SMBP first, one after the other	189	0.59/0.64		0.62/0.67	0.62/0.67		0.66/0.70
Verberk (2006)	7 days (3 readings in the morning and 3 readings in the evening, with the first of each triplicate discarded)		Daytime ABPM Timing vs. SMBP unclear	216				0.63/0.65	0.58/0.60	0.65/0.66
		24- Tin u	24-hour ABPM Timing vs. SMBP unclear	216				0.69/0.69	0.64/0.62	0.70/0.69
Study.			Correlation coefficient between home systolic/ diastolic BP and th	Correlat	tion coefficient b	letween home sy	stolic/ diastolic	Correlation coefficient between home systolic/ diastolic BP and the comparator measurement	arator measurem	nent
(first author; publication date)	Measurement schedule	Comparator	2 7	2	3 4	5678	9 10 11	12 16	20	24 28
Johannson (2010)	7 days (2 readings in the 2 morning and 2 readings T in the evening on each day)	24-hour ABPM Timing in relationship to SMBP unclear	464		0.84/0.82	0.87/0.84		0.88/0.85 0.88/0.86	0.89/0.86	0.89/0.87 0.89/0.87
Stergiou (1998)	3 work days per week for D 2 weeks (2 readings S in the morning and 2 readings in the evening on each day)	Daytime ABPM Some ABPM first and some SMBP first, one after the other	189 0.59/0.64	0.62/0.67	0.66/0.70	0.68/0.73		0.68/0.75 0.70/0.77	0.71/0.78	0.71/0.79
Verberk (2006)		Daytime ABPM Timing vs. SMBP unclear	216 0	0.53/0.62	2 0.53/0.59	0.57/0.61		0.60/0.63 0.62/0.63	0.63/0.66	0.64/0.66 0.65/0.66
	evening, with the `first of each triplicate 2, discarded)	24-hour ABPM Timing vs. SMBP unclear	216 0	0.57/0.65	0.57/0.61	0.62/0.63		0.66/0.66 0.68/0.66	0.68/0.66 0.69/0.69 0.6	0.69/0.69 0.70/0.69

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American Journal of Hypertension 13

inform individuals where more than 3 days of measurements are appropriate.

These data hold for both diagnosis and ongoing management. There are theoretical reasons (peaks and troughs of medication for example) that support recommendations for morning and evening readings.²³ In terms of diagnosis, the prognostic studies did not suggest any particular difference in time of measurement and neither were differences in correlation seen dependent on time of day of monitoring, perhaps suggesting that such considerations are not paramount.

On the basis of the evidence we have synthesized, a pragmatic revision of current guidelines for self-monitoring would be that measurement of BP should be undertaken for 3 days, whether for diagnostic purposes or when monitoring the effect of treatment change, unless mean blood pressure after 3 days is close to a treatment or diagnostic threshold when longer schedules—perhaps a further 3 days of monitoring—bring small increases in prognostic power. Precise timings of measurements within these days and the precise days of measurement are less important and might be varied to suit individual circumstances. There remains a need for more and higher quality research, particularly prognostic studies in diverse populations, involving comparison of different regimes of measurement across multiple studies.

DATA AVAILABILITY

Dataset available from the corresponding author at j.a.hodgkinson@bham.ac.uk. The dataset includes only anonymized material already in the public domain.

SUPPLEMENTARY DATA

Supplementary data are available at *American Journal of Hypertension* online.

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DISCLOSURE

All authors have completed the ICMJE uniform disclosure form at www.icmje.org/coi_disclosure.pdf (available on request from the corresponding author) and declare no support from any organization for the submitted work, no financial relationships with any organizations that might have an interest in the submitted work in the previous 3 years, and no other relationships or activities that could appear to have influenced the submitted work.

REFERENCES

- 1. Lim SS, Vos T, Flaxman AD, Danaei G, Shibuya K, Adair-Rohani H, Amann M, Anderson HR, Andrews KG, Aryee M, Atkinson C, Bacchus LJ, Bahalim AN, Balakrishnan K, Balmes J, Barker-Collo S, Baxter A, Bell ML, Blore JD, Blyth F, Bonner C, Borges G, Bourne R, Boussinesq M, Brauer M, Brooks P, Bruce NG, Brunekreef B, Bryan-Hancock C, Bucello C, Buchbinder R, Bull F, Burnett RT, Byers TE, Calabria B, Carapetis J, Carnahan E, Chafe Z, Charlson F, Chen H, Chen JS, Cheng AT, Child JC, Cohen A, Colson KE, Cowie BC, Darby S, Darling S, Davis A, Degenhardt L, Dentener F, Des Jarlais DC, Devries K, Dherani M, Ding EL, Dorsey ER, Driscoll T, Edmond K, Ali SE, Engell RE, Erwin PJ, Fahimi S, Falder G, Farzadfar F, Ferrari A, Finucane MM, Flaxman S, Fowkes FG, Freedman G, Freeman MK, Gakidou E, Ghosh S, Giovannucci E, Gmel G, Graham K, Grainger R, Grant B, Gunnell D, Gutierrez HR, Hall W, Hoek HW, Hogan A, Hosgood HD, III, Hoy D, Hu H, Hubbell BJ, Hutchings SJ, Ibeanusi SE, Jacklyn GL, Jasrasaria R, Jonas JB, Kan H, Kanis JA, Kassebaum N, Kawakami N, Khang YH, Khatibzadeh S, Khoo JP, Kok C, Laden F, Lalloo R, Lan Q, Lathlean T, Leasher JL, Leigh J, Li Y, Lin JK, Lipshultz SE, London S, Lozano R, Lu Y, Mak J, Malekzadeh R, Mallinger L, Marcenes W, March L, Marks R, Martin R, McGale P, McGrath J, Mehta S, Mensah GA, Merriman TR, Micha R, Michaud C, Mishra V, Mohd Hanafiah K, Mokdad AA, Morawska L, Mozaffarian D, Murphy T, Naghavi M, Neal B, Nelson PK, Nolla JM, Norman R, Olives C, Omer SB, Orchard J, Osborne R, Ostro B, Page A, Pandey KD, Parry CD, Passmore E, Patra J, Pearce N, Pelizzari PM, Petzold M, Phillips MR, Pope D, Pope CA, III, Powles J, Rao M, Razavi H, Rehfuess EA, Rehm JT, Ritz B, Rivara FP, Roberts T, Robinson C, Rodriguez-Portales JA, Romieu I, Room R, Rosenfeld LC, Roy A, Rushton L, Salomon JA, Sampson U, Sanchez-Riera L, Sanman E, Sapkota A, Seedat S, Shi P, Shield K, Shivakoti R, Singh GM, Sleet DA, Smith E, Smith KR, Stapelberg NJ, Steenland K, Stöckl H, Stovner LJ, Straif K, Straney L, Thurston GD, Tran JH, Van Dingenen R, van Donkelaar A, Veerman JL, Vijayakumar L, Weintraub R, Weissman MM, White RA, Whiteford H, Wiersma ST, Wilkinson JD, Williams HC, Williams W, Wilson N, Woolf AD, Yip P, Zielinski JM, Lopez AD, Murray CJ, Ezzati M, AlMazroa MA, Memish ZA. A comparative risk assessment of burden of disease and injury attributable to 67 risk factors and risk factor clusters in 21 regions, 1990-2010: a systematic analysis for the Global Burden of Disease Study 2010. Lancet 2012; 380:2224-2260.
- Excellence NIfC. Hypertension: Management of Hypertension in Adults in Primary Care. http://www.nice.org.uk/guidance/CG127/ NICEGuidance; 2011. Report No: Clinical Guideline 127.
- 3. Mancia G, Fagard R, Narkiewicz K, Redón J, Zanchetti A, Böhm M, Christiaens T, Cifkova R, De Backer G, Dominiczak A, Galderisi M, Grobbee DE, Jaarsma T, Kirchhof P, Kjeldsen SE, Laurent S, Manolis AJ, Nilsson PM, Ruilope LM, Schmieder RE, Sirnes PA, Sleight P, Viigimaa M, Waeber B, Zannad F; Task Force Members. 2013 ESH/ESC Guidelines for the management of arterial hypertension: the Task Force for the management of arterial hypertension of the European Society of Hypertension (ESH) and of the European Society of Cardiology (ESC). J Hypertens 2013; 31:1281–1357.
- Pickering TG, Hall JE, Appel LJ, Falkner BE, Graves JW, Hill MN, Jones DH, Kurtz T, Sheps SG, Roccella EJ; Council on High Blood

- Hodgkinson J, Mant J, Martin U, Guo B, Hobbs FD, Deeks JJ, Heneghan C, Roberts N, McManus RJ. Relative effectiveness of clinic and home blood pressure monitoring compared with ambulatory blood pressure monitoring in diagnosis of hypertension: systematic review. *BMJ* 2011; 342:d3621.
- McManus RJ, Glasziou P, Hayen A, Mant J, Padfield P, Potter J, Bray EP, Mant D. Blood pressure self monitoring: questions and answers from a national conference. *BMJ* 2008; 337:a2732.
- Ward AM, Takahashi O, Stevens R, Heneghan C. Home measurement of blood pressure and cardiovascular disease: systematic review and meta-analysis of prospective studies. *J Hypertens* 2012; 30:449–456.
- 8. Shimamoto K, Ando K, Fujita T, Hasebe N, Higaki J, Horiuchi M, Imai Y, Imaizumi T, Ishimitsu T, Ito M, Ito S, Itoh H, Iwao H, Kai H, Kario K, Kashihara N, Kawano Y, Kim-Mitsuyama S, Kimura G, Kohara K, Komuro I, Kumagai H, Matsuura H, Miura K, Morishita R, Naruse M, Node K, Ohya Y, Rakugi H, Saito I, Saitoh S, Shimada K, Shimosawa T, Suzuki H, Tamura K, Tanahashi N, Tsuchihashi T, Uchiyama M, Ueda S, Umemura S, on behalf of The Japanese Society of Hypertension Committee for Guidelines for the Management of Hypertension. (JSH 2014). *Hypertens Res* 2014; 37:253–392.
- Pickering TG, Miller NH, Ogedegbe G, Krakoff LR, Artinian NT, Goff D; American Heart Association; American Society of Hypertension; Preventive Cardiovascular Nurses Association. Call to action on use and reimbursement for home blood pressure monitoring: a joint scientific statement from the American Heart Association, American Society of Hypertension, and Preventive Cardiovascular Nurses Association. *Hypertension* 2008; 52:10–29.
- EPHPP. Effective Public Health Practice Project Quality Assessment Tool (EPHPP). https://merst.ca/wp-content/uploads/2018/02/qualityassessment-tool_2010.pdf
- 11. Downs SH, Black N. The feasibility of creating a checklist for the assessment of the methodological quality both of randomised and non-randomised studies of health care interventions. *J Epidemiol Community Health* 1998; 52:377–384.

- 12. Whiting PF, Rutjes AW, Westwood ME, Mallett S, Deeks JJ, Reitsma JB, Leeflang MM, Sterne JA, Bossuyt PM; QUADAS-2 Group. QUADAS-2: a revised tool for the quality assessment of diagnostic accuracy studies. *Ann Intern Med* 2011; 155:529–536.
- O'Brien E, Amoore JN, Asmar R, Atkins N, Beilin L, Bouchier-Hayes D. dabl Educational Trust [cited 2018]. http://www.dableducational.org
- British and Irish Hypertension Society. Validated BP Monitors for Home Use [cited 2018]. https://bihsoc.org/bp-monitors/for-home-use/
- Hughes MD. Regression dilution in the proportional hazards model. *Biometrics* 1993; 49:1056–1066.
- 16. Niu K, Hozawa A, Awata S, Guo H, Kuriyama S, Seki T, Ohmori-Matsuda K, Nakaya N, Ebihara S, Wang Y, Tsuji I, Nagatomi R. Home blood pressure is associated with depressive symptoms in an elderly population aged 70 years and over: a population-based, cross-sectional analysis. *Hypertens Res* 2008; 31:409–416.
- Nakagawa H, Niu K, Hozawa A, Ikeda Y, Kaiho Y, Ohmori-Matsuda K, Nakaya N, Kuriyama S, Ebihara S, Nagatomi R, Tsuji I, Arai Y. Impact of nocturia on bone fracture and mortality in older individuals: a Japanese longitudinal cohort study. *J Urol* 2010; 184:1413–1418.
- Stergiou GS, Baibas NM, Kalogeropoulos PG. Cardiovascular risk prediction based on home blood pressure measurement: the Didima study. J Hypertens 2007; 25:1590–1596.
- Verberk ŴJ, Kroon AA, Kessels AG, de Leeuw PW. Home blood pressure measurement: a systematic review. J Am Coll Cardiol 2005; 46:743–751.
- Niiranen TJ, Asayama K, Thijs L, Johansson JK, Hara A, Hozawa A, Tsuji I, Ohkubo T, Jula AM, Imai Y, Staessen JA; IDHOCO Investigators. Optimal number of days for home blood pressure measurement. *Am J Hypertens* 2015; 28:595–603.
- 21. Grant S, Hodgkinson JA, Milner SL, Martin U, Tompson A, Hobbs FR, Mant J, McManus RJ, Greenfield SM. Patients' and clinicians' views on the optimum schedules for self-monitoring of blood pressure: a qualitative focus group and interview study. *Br J Gen Pract* 2016; 66:e819–e830.
- van der Hoeven NV, van den Born BJ, Cammenga M, van Montfrans GA. Poor adherence to home blood pressure measurement schedule. J Hypertens 2009; 27:275–279.
- 23. Imai Y, Nishiyama A, Sekino M, Aihara A, Kikuya M, Ohkubo T, Matsubara M, Hozawa A, Tsuji I, Ito S, Satoh H, Nagai K, Hisamichi S. Characteristics of blood pressure measured at home in the morning and in the evening: the Ohasama study. J Hypertens 1999; 17:889–898.