Interarm Difference in Systolic Blood Pressure in Different Ethnic Groups and Relationship to the “White Coat Effect”: A Cross-Sectional Study

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BACKGROUND
Interarm differences (IADs) ≥10 mm Hg in systolic blood pressure (BP) are associated with greater incidence of cardiovascular disease. The effect of ethnicity and the white coat effect (WCE) on significant systolic IADs (ssIADs) are not well understood.

METHODS
Differences in BP by ethnicity for different methods of BP measurement were examined in 770 people (300 White British, 241 South Asian, 229 African-Caribbean). Repeated clinic measurements were obtained simultaneously in the right and left arm using 2 BPTru monitors and comparisons made between the first reading, mean of second and third and mean of second to sixth readings for patients with, and without known hypertension. All patients had ambulatory BP monitoring (ABPM). WCE was defined as systolic clinic BP ≥10 mm Hg higher than daytime ABPM.

RESULTS
No significant differences were seen in the prevalence of ssIAD between ethnicities whichever combinations of BP measurement were used and regardless of hypertensive status. ssIADs fell between the 1st measurement (161, 22%), 2nd/3rd (113, 16%), and 2nd-6th (78, 11%) (1st vs. 2nd/3rd and 2nd-6th, P < 0.001). Hypertensives with a WCE were more likely to have ssIADs on 1st, (odds ratio [OR] 1.73 (95% confidence interval 1.04–2.86); 2nd/3rd, (OR 3.05 (1.68–5.53)); and 2nd-6th measurements, (OR 2.58 (1.22–5.44). Nonhypertensive participants with a WCE were more likely to have a ssIAD on their first measurement (OR 3.82 (1.77 to −8.25) only.

CONCLUSIONS
ssIAD prevalence does not vary with ethnicity regardless of hypertensive status but is affected by the number of readings, suggesting the influence of WCE. Multiple readings should be used to confirm ssIADs.

Keywords: ambulatory blood pressure monitoring; blood pressure; ethnic group; hypertension; interarm blood pressure differences; simultaneous blood pressure measurement method; white coat effect.

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Differences in systolic blood pressure (BP) between right and left arms ≥10 mm Hg independently predict increased risk of cardiovascular events,1 subclavian stenosis,2 and cardiovascular or all-cause mortality.3 Measuring BP in both arms is a simple intervention that identifies the higher reading arm, which should be used for future hypertension management, and may identify patients needing investigation, or intensification of cardiovascular risk management.4 Prevalence of
patients with a significant systolic interarm difference (ssIAD) varies according to population. A U.S. systematic review, found the prevalence of ssIADs was 7.5% in primary care, 9% for hospital outpatients, and 12.1% for hospital inpatients. A recent study using Framingham data found the prevalence of ssIADs in a community sample, free from cardiovascular disease at baseline, was 9.4%. A UK systematic review considered prevalence of ssIADs in relation to clinical predictors and found 11.2% in hypertensives, 7.4% in people with diabetes and 3.6% in the general population. Characteristics associated with ssIADs are age, diabetes, higher systolic BP and increased body mass index (BMI).

Some ethnic groups are at higher cardiovascular risk. South Asians and African-Caribbeans have a higher incidence of diabetes than White Europeans, and African-Caribbeans have a greater prevalence of hypertension and stroke. Ethnicity may also affect the prevalence of ssIADs. Countries in the Far East appear to have a lower prevalence of patients with ssIADs than Western populations.

To our knowledge, there has only been one study so far to evaluate whether prevalence of ssIADs are age, diabetes, race, and African-Caribbean ethnicity, hypertensive status or other participant characteristic and additionally to investigate any association of prevalence with the number of readings, or the presence of a white coat effect (WCE) or white coat hypertension (WCH).

**METHODS**

Blood pressure monitoring in different ethnic groups (BP-ETH), was a primary care based observational study conducted between June 2010 and December 2012. The detailed methods have been published previously and are outlined below.

**Population**

Participants were aged 40–74 years, purposefully recruited from 3 ethnic groups: White British (WB), South Asian (SA), and African-Caribbean (AC). Ethnicity was self-defined using standard UK criteria. Respondents attended 3 research clinics at their own practice. Twenty-eight practices were recruited from the Primary-Care-Research-Network, Central England, United Kingdom, chosen to represent the required range of ethnicities. Around 40 participants were recruited from each practice, both with (HT) and without hypertension (NHT), as defined by a clinical code in the patient’s record. People unable to consent, belonging to a different ethnic group or whose general practitioner felt they were unable to take part were excluded.

**Procedure**

Following at least 5 minutes rest, sitting BP measurements were taken by a research nurse using 2 BPTru monitors (BPTru Medical Devices BPM-200), set to take 6 readings at 1-minute intervals and used simultaneously on both arms. Monitors were calibrated independently to the same standard at the start of the study and the BPTru device performs an autocalibration upon activation to maintain accuracy. The research nurse remained in the room with the patient while the readings were being taken. One BPTru monitor was used consistently for the right arm and the second used for the left. This is considered to be the most accurate way of establishing an IAD.

Participants were fitted with an ambulatory monitor (Spacelabs 90217-1Q) either on the first or second clinic visit. Ambulatory BP monitoring (ABPM) readings were recorded at half hourly intervals during the day and hourly overnight for a total of 24 hours. Participants’ nondominant arm was used unless systolic BP was ≥20 mm Hg between the right and left arm on the first reading, which included 32 patients. In the case of these patients, the arm with the higher reading was used.

**Analysis**

ssIAD for the first measurement was defined as an absolute difference ≥10 mm Hg between the right and left arm. SSIIADs were further defined using absolute ssIADs of the 1st reading alone, mean of the 2nd/3rd and mean of the 2nd to 6th readings.

There is no accepted definition of the WCE or WCH therefore different analyses were undertaken based on the literature using the following systolic BP definitions:

1. **White coat effect (1st BP measurement):**
   a. (1st clinic measurement–mean daytime ABPM) ≥10 mm Hg
2. **White coat effect (mean of 2nd/3rd BP measurements)**
   b. (Mean of 2nd/3rd clinic measurement–mean daytime ABPM) ≥10 mm Hg
3. **White coat effect (2nd–6th BP measurement)**
   c. (Mean of 2nd–6th clinic measurement–mean daytime ABPM) ≥10 mm Hg

Trends between the ethnic groups for prevalence of ssIADs were tested using the Extended Mantel-Haenszel chi-squared test for linear trend.

Univariable analyses were conducted between ssIADs and baseline characteristics stratified by diagnosed hypertension (HT) and not known to be hypertensive (NHT) patients. Chi-squared test was used when comparing binary/categorical variables. The 2 sample t-test was used when comparing the mean difference between 2 groups. Nonparametric tests were used for skewed variables such as BMI.

A multivariate logistic regression model was also used where we adjusted for the following variables: ethnicity,
diabetes, age, gender, CHD, log(BMI), mean BP, medication, smoking, and deprivation score.

Differences in the proportion of patients with an absolute ssIAD, dependent on the number of measurements, were investigated using a Cochran Q test. All BP comparisons were for measurements in the same arm.

The same analysis was conducted checking for any differences in the prevalence of the WCE across different combinations of measurements. Post-hoc analyses were conducted using the McNemar test. Association between ssIADs and the corresponding WCE was examined by the use of logistic regression models adjusted for baseline characteristics.

Sensitivity analyses were carried out to check the effect on WCE when using differences between clinic systolic BP and systolic ABPM of 5 and 15 mm Hg.

Ethics and research governance approval

Ethical approval for the BP-ETH study was gained from the Black Country Research Ethics Committee, West Midlands, United Kingdom (Ref 09/H1202//114).

RESULTS

Baseline demographic data

Seven hundred and seventy people participated in BP-ETH of whom 300 were WB, 229 were AC, and 241 were SA (Supplementary Table 1a—Appendix). There were no significant differences in baseline characteristics between WB, AC, and SA groups, or between HT and NHT participants, although more HT participants (481) were recruited than NHT (289).

Prevalence of IADs by ethnicity and hypertensive status

BP measurements in the right and left arms were available for 750 of the 770 participants who were grouped into HT and NHT and then subdivided into participants with a ssIAD or not. There was no systematic difference in the systolic BP of the right arm vs. that of the left (Supplementary Table 2a—Appendix); therefore, the following ssIADs are expressed as an absolute difference.

The overall prevalence of ssIADs for HT participants was 61/469 (13.0%) and 26/281 (9.3%) for NHT participants (Table 1). Overall prevalence of ssIADs in the HT group was higher but this difference was not found to be significant (Table 1). Association between ssIAD and ethnic group was not significant for the HT or NHT group (Table 1). There were no significant differences in the prevalence of ssIAD by ethnicity in the HT and NHT groups whichever combination of measurements was used (1\(^{st}\), mean of 2\(^{nd}\)/3\(^{rd}\), or mean of 2\(^{nd}\)-6\(^{th}\)). Therefore only the ssIAD using the mean of the 2\(^{nd}\)-6\(^{th}\) measurement is presented here as prevalence dropped the more measurements were used and this mean represented the nadir. A multivariate logistic regression model was used where we adjusted for diabetes, age, gender, CHD, log(BMI), deprivation score, and mean BP. Ethnicity remained nonsignificant in both HT and NHT patients (Table 3).

### Characteristics of participants with and without an IAD

Table 2 shows characteristics of participants with and without a ssIAD by hypertensive status. As the prevalence of ssIADs was not significantly associated with ethnicity, the 3 ethnic groups are combined for analysis of the remaining characteristics. Significantly higher mean BP was associated with a ssIAD for both HT (140.1 mm Hg [ssIAD] vs. 131.6 mm Hg [no ssIAD], \(P < 0.001\)) and NHT groups (132.0 mm Hg [ssIAD] vs. 124.7 mm Hg [no ssIAD], \(P = 0.031\)) (Table 2). The multivariate analysis also showed a significantly higher BP was associated with a ssIAD in both the HT and NHT groups (Table 3). Significantly higher daytime ABPM was associated with a ssIAD in the HT but not the NHT group.

As BMI was a skewed variable, this is reported as a median value with an interquartile range using the nonparametric test Wilcoxon rank sum for the \(P\) value (Table 2). Participants with a ssIAD had a significantly higher BMI in the HT group (31.0 kg/m\(^2\) vs. 28.7 kg/m\(^2\), \(P = 0.025\)); however, this significance disappeared in the multivariate analysis (Table 3). Similarly in the NHT group, where ssIAD was actually associated with a lower BMI (27.3 kg/m\(^2\) vs. 29.4 kg/m\(^2\), \(P = 0.025\)) (Table 2), the BMI no longer remained significant in the multivariate analysis (Table 3). Multivariate analysis showed that NHT participants with a ssIAD were significantly more likely to smoke, which may have affected the univariate analysis of BMI.

Difference in age between HT participants with a ssIAD and HT participants without a ssIAD was of borderline significance (58.4 vs. 60.8, \(P = 0.053\)). This effect was not seen

### Table 1. Incidence of ssIADs by hypertensive status and ethnic group

|                  | Known hypertensive (HT) | Not known to be hypertensive (NHT) |
|------------------|-------------------------|-----------------------------------|
|                  | \(\geq 10\) mm Hg (\(n = 61\)) | <10 mm Hg (\(n = 408\)) | \(P\) value | \(\geq 10\) mm Hg (\(n = 26\)) | <10 mm Hg (\(n = 255\)) | \(P\) value |
| Overall, No. (%) | 61 (13.0) | 408 (87.0) | | 26 (9.3) | 255 (90.7) | |
| WB, No. (%)      | 17 (40) | 162 (28) | 0.188 | 10 (38) | 101 (40) | 0.892 |
| SA, No. (%)      | 20 (29) | 120 (33) | | 10 (38) | 87 (34) | |
| AC, No. (%)      | 24 (31) | 126 (39) | | 6 (23) | 67 (26) | |

Abbreviations: AC, African-Caribbean; BP, blood pressure; SA, South Asian; ssIAD, Systolic Interarm Differences; WB, White British.
in the NHT group and the effect was no longer significant when other variables were accounted for in the multivariate analysis (Table 3). There were no other significant differences between participants with and without ssIAD (Table 2).

Table 2. Showing demographic and health differences between those patients with an interarm BP difference and those without for the HT and NHT groups

| Variable                      | Known hypertensive (HT) | Not known to be hypertensive (NHT) | \( P \) value |
|-------------------------------|-------------------------|-----------------------------------|---------------|
|                               | \( \geq 10 \) mm Hg (\( n = 61 \)) | \( <10 \) mm Hg (\( n = 408 \)) |               |
| Age (years)                   | 58.4 (8.9)              | 60.8 (8.8)                        | 0.053         |
| Men                           | 26 (43%)                | 208 (51%)                         | 0.223         |
| Mean blood Pressure\(^a\)     | (\( n = 52 \))          | (\( n = 386 \))                   | <0.001        |
| SD                            | 143.7 (19.6)            | 134.1 (17.6)                      |               |
| BMI\(^b\) (kg/m\(^2\))       | 31.0 (26.7–34.3)        | 28.7 (25.5–32.1)                  | 0.0254        |
| Coronary heart disease        | 5 (8%)                  | 53 (13%)                          | 0.403         |
| Diabetes                      | 16 (27%)                | 94 (23%)                          | 0.536         |
| Mean deprivation score\(^c\) | (\( n = 60 \))          | (\( n = 384 \))                   | 0.572         |
| SD                            | 44.1 (17.55)            | 42.8 (17.58)                      |               |
| Mean daytime ABPM\(^a\)      | (\( n = 47 \))          | (\( n = 341 \))                   | 0.003         |
| SD                            | 139.5 (15.8)            | 132.8 (14.3)                      |               |
| Taking antihypertensive       | 58 (95%)                | 396 (97%)                         | 0.308         |
| Smoking (Y)                   | 6 (10%)                 | 60 (15%)                          | 0.207         |

Abbreviations: ABPM, ambulatory blood pressure monitoring BP, blood pressure.
\(^a\)Number of participants unless otherwise stated.
\(^b\)mm Hg.
\(^c\)BMI, body mass index, reported as a median value with (interquartile range).
\(^d\)Index of multiple deprivation 2007.

Table 3. Multivariable analysis showing the likelihood of a ssIAD for the HT and NHT groups

| Variables                        | Known hypertensive (HT) (\( n = 437 \)) (OR) | Not known to be hypertensive (NHT) (\( n = 262 \)) (OR) | \( P \) value |
|----------------------------------|----------------------------------------------|---------------------------------------------------------|---------------|
| WB\(^a\)                         | 1                                            | 1                                                       |               |
| SA                               | 1.9 (0.851–4.333)                            | 1.2 (0.386–3.916)                                        |               |
| AC                               | 1.7 (0.777–3.919)                            | 0.8 (0.226–2.963)                                        |               |
| Diabetes                         | 1.2 (0.597–2.483)                            | 0.5 (0.0524–5.495)                                       |               |
| Age                              | 1.0 (0.940–1.011)                            | 1.0 (0.936–1.058)                                        |               |
| Sex (Men)                        | 0.8 (0.446–1.527)                            | 0.8 (0.324–2.128)                                        |               |
| CHD                              | 0.6 (0.177–2.169)                            | 1.4 (0.0663–29.91)                                       |               |
| BMI\(^b\) (log transformed)      | 2.6 (0.580–11.46)                            | 0.2 (0.0141–3.170)                                       |               |
| Mean blood pressure              | 1.03*** (1.010–1.043)                         | 1.03** (1.006–1.064)                                     |               |
| On medication                    | 0.8 (0.158–4.160)                            | 0.8 (0.0410–15.17)                                       |               |
| Smoker                           | 0.9 (0.333–2.184)                            | 5.4*** (1.887–15.21)                                     |               |
| IMD\(^c\)                        | 1.0 (0.988–1.004)                            | 1.0 (1.000–1.003)                                        |               |

Abbreviations: AC, African-Caribbean; CHD, coronary heart disease; OR, odds ratio; SA, South Asian; ssIAD, significant systolic interarm differences; WB, White British.
\(^a\)Reference category ***\( P < 0.01 \)**\( P < 0.05 \).
\(^b\)Body mass index.
\(^c\)Index of multiple deprivation 2007.

Changes in interarm BP differences between the 1\(^{st}\) reading, 2\(^{nd}\) and 3\(^{rd}\) reading, and 2–6\(^{th}\) reading

Out of 469 HT participants, 449 had all 6 BP readings available on both arms. The number of participants with a
ssIAD fell as more pairs of readings were included in the calculation of mean BP (Supplementary Table 3a—Appendix): 101 (22%) for the 1st measurement, 69 (15%) for the mean of the 2nd–3rd measurements, and 54 (12%) for the mean of the 2nd–6th measurements. Post-hoc analysis revealed a significant difference \( (P < 0.001) \) between the number of participants with a ssIAD on the 1st pair of readings vs. the number with a ssIAD on the mean of the 2nd–3rd and the 2nd–6th pairs of readings, with a smaller nonsignificant difference between the latter 2 measurement methods (Supplementary Table 3a—Appendix). NHT (271/281) patients showed a similar pattern (Supplementary Table 3a—Appendix).

This effect was mirrored by the decline in the prevalence of a WCE; numbers of HT participants with a WCE on the 1st measurement was 128 (35%), 89 (25%) for the mean of the 2nd–3rd, and 66 (18%) for 2nd–6th BP readings (1st vs. 2nd/3rd \( P < 0.001; 1 \text{st} \) vs. 2nd–6th reading \( P < 0.001 \) (Supplementary Table 4a—Appendix). This decline was also seen in the NHT participants (Supplementary Table 4a—Appendix).

### Association of the WCE and interarm BP difference

The relationship between the WCE and ssIAD was investigated using a logistic regression model and adjusted for baseline characteristics. Resulting odds ratios show that in the HT group there was a significant association between the WCE and ssIAD for the 1st measurement, 2nd–3rd, and 2nd–6th measurement (Table 4).

In the NHT group, the odds ratio was significant for the association between WCH and ssIAD for the 1st measurement \( (4.06, 95\% \text{ confidence interval}, 1.83–9.00 \) (Table 4) but no significant differences were found in the association of WCH and ssIAD between the mean of the 2nd–3rd and the 2nd–6th readings (Table 4).

Sensitivity analyses using definitions of a 5 mm Hg and 15 mm Hg difference between clinic systolic BP and the systolic mean daytime ABPM showed a similar pattern of results (Supplementary Tables 5a and 6a—Appendix).

### DISCUSSION

#### Summary

In this community-based study, ethnicity had no significant impact on the prevalence of ssIAD. The HT group had a greater prevalence of ssIADs overall but the difference between the HT and the NHT groups was not significant. However, the prevalence of ssIADs significantly increased with increasing BP regardless of hypertensive status. NHT participants with a ssIAD were more likely to be a smoker, but there did not appear to be the same association for BMI and age. There was no systematic difference in systolic BP between the right and the left arm, which was in keeping with the results of a recent meta-analysis.

Calculating ssIAD with increasing numbers of repeated measurements significantly reduced the prevalence of a ssIAD and this appeared to be associated with the WCE, especially for HT participants. For NHT participants, the association between a ssIAD and WCH was present on the first measurement but not thereafter.

### Strengths and Limitations

The main strength of this study was that it was large, community-based, and recruited approximately equal numbers of patients from each of 3 ethnic groups. It included patients with a diagnosis of hypertension, those with no known diagnosis and did not exclude patients with comorbidities. Measurements were taken in a Primary Care setting, which is where most office BP is taken and provides the most generalizable comparison of the WCE. The study used 6 BP measurements, allowing for analysis of ssIADs over several consecutive readings, and measurements were taken simultaneously with a validated monitor which is widely acknowledged as the most accurate way to measure IAD. There is a theoretical bias arising from the use of different monitors on each arm, which could not be helped as the BPTTru monitor takes readings automatically at 1-minute intervals precluding switching of monitors between arms. However, the monitors were all calibrated prior to the start of the study and auto-calibrate each time when activated therefore reducing the likelihood of monitors on different arms being in a different state of calibration. The number of patients with ssIADs was relatively small which affected power to assess associations and differences between groups. Only systolic BP was assessed therefore any associated or independent effects on IAD between diastolic BP are unknown.

The current study did not include Far Eastern ethnicities such as Chinese, who may have significantly lower incidence of ssIADs than Western groups. Despite the range of ethnicities, there was only 1 area of the UK studied, which does not take into account potential effects different environments may have.

The population here came from more deprived areas in comparison with the rest of the UK. While we do not know what association deprivation would have with the prevalence

**Table 4.** ORs for ssIADs when the WCE is present in HT and NHT groups

|                                | ORs for ssIAD when WCE is present in HT group | ORs for ssIAD when WCE is present in NHT group |
|--------------------------------|---------------------------------------------|-----------------------------------------------|
| First pair of readings         | 2.12 (95% CI: 1.24–3.62; \( P = 0.006 \), \( n = 370 \)) | 4.06 (95% CI: 1.83–9.00; \( P = 0.001 \), \( n = 220 \)) |
| Mean of the 2nd and 3rd pair of readings | 3.69 (95% CI: 1.96–6.93; \( P < 0.001 \), \( n = 373 \)) | 1.07 (95% CI: 0.41–2.79; \( P = 0.888 \), \( n = 226 \)) |
| Mean of the 2nd–6th pair of readings | 3.48 (95% CI: 1.56; 7.71; \( P = 0.002 \), \( n = 364 \)) | 1.68 (95% CI: 0.451–6.32; \( P = 0.436 \), \( n = 219 \)) |

Logistic regression model was used and adjusted for gender, ethnicity, logarithm of BMI, age, CHD, daytime ABPM, diabetes, smoking, medication, and IMID. Abbreviations: CI, confidence interval; HT, known hypertensive; IMID, index of multiple deprivation; NHT, not known to be hypertensive; OR, odds ratio; ssIAD, significant systolic interarm differences; WCE, white coat effect.
of ssIADs, hypertension is more prevalent in deprived settings so a concomitant trend to increased prevalence of ssIADs with deprivation might be predicted. In this study, the definition of a clinically significant systolic IAD was defined as ≥10 mm Hg. Current European hypertension guidelines suggest a ssIAD >10 mm Hg between arms carries an increase in cardiovascular risk and although the UK guidelines suggest a difference in systolic BP ≥20 mm Hg is indicative of vascular disease, they only specify a ssIAD <10 mm Hg as normal. Although a ssIAD ≥10 mm Hg may show less specificity for peripheral artery disease than that of 15 mm Hg, it is more sensitive and, in addition, it is in keeping with the definition used in many current studies. Using this definition meant that our work could be compared to the current literature more easily and was clinically relevant.

As there is no accepted definition of the WCE or WCH, a pragmatic definition for the study was developed using an arbitrary level of the clinic-ambulatory difference (≥10 mm Hg). Differences between clinic and ambulatory BP are potentially subject to bias, principally from variation in the clinic BP due to operator error, hypertensive status and activities, and environment of the patient. However, given the controlled nature of the research measurements, such errors would be expected to be minimized in this study.

Comparison with existing literature

The prevalence of ssIADs in the HT and NHT groups was 13.0% and 9.3%, respectively. In the NHT group, this appeared to be high for a general population.

A recent systematic review, analysing prevalence of ssIADs from 16 studies, found a prevalence of 3.6%. However, the population group included 2 studies of Far Eastern origin which had a much lower prevalence than studies of Western origin and therefore reduced the pooled prevalence. There was a higher prevalence in a hypertensive population and when studies causing statistical heterogeneity were removed, the prevalence of ssIADs in a Western, hypertensive population was 13.3%, almost identical to the prevalence found here. Similar to the current study, the multiethnic study of atherosclerosis (MESA) also reported no significant differences in ssIADs between African American and White non-Hispanic groups. However, there was a significantly lower prevalence of ssIADs in Hispanic and Chinese Americans suggesting that Hispanic and Far Eastern ethnicities may be less predisposed to ssIADs than Western populations.

A higher BMI has been previously reported as being associated with a ssIAD and is likely to relate to patients with ssIADs having a higher cardiovascular risk. However, another study found that age was the only significant predictor for ssIADs. Neither age nor BMI was found to have a significant association here with a ssIAD in the HT or NHT group. However, BMI was higher in the HT group so this may have been compounded by the fact that the number of patients with a ssIAD was small. The NHT group had a significant association between smoking and ssIAD, which is in keeping with the link between ssIADs and a greater cardiovascular risk. However, the link between smoking and ssIADs is varied with some studies showing some association but others showing no significance. The number of participants in this study who smoked was very small and is likely to affect the significance of any associations here.

Results here are in agreement with that found in the Framingham Heart Study, a study in a Japanese population and a recent large meta-analysis of 16 IAD studies showing that those patients with a ≥10 mm Hg difference between arms had significantly higher systolic BP compared to those without. This is likely to be an effect of the absolute IAD increasing as absolute BP increases and may in part explain why patients with a ssIAD appear to be at higher risk of a cardiovascular event.

Systolic BP decreased over the 6 clinic measurements, and the prevalence of ssIAD followed the same pattern. This pattern has been seen in other studies and a study by Martin et al. suggested that the effect may be associated with the WCE, although they used sequential measurements to estimate IAD which can overestimate its prevalence.

A significant association was found here between WCH and ssIAD on the first reading for participants in the NHT group and between the WCE and ssIADs for all BP measurements used in the HT patients.

For HT patients, the strongest association between the WCE and ssIAD was using the mean of 2nd/3rd readings which may be explained by HT participants having a stronger, more persistent WCE than those with no diagnosis. There is closer agreement between clinic and ambulatory BP when a patient's BP is closer to normal levels. This suggests that the increase in prevalence of ssIADs in response to higher BP levels may be linked to the WCE.

Implications for practice

There appears to be little difference in the prevalence of ssIADs between SA and AC cohorts compared to a WB population. However, those with higher mean clinic BP were more likely to have a ssIAD regardless of hypertensive status. European guidelines recommend simultaneous measurement to exclude clinical ssIADs. A much greater effect was seen in terms of the number of measurements used; hence, health professionals should not rely on single BP measurements to identify ssIADs.

This study and others have shown that the prevalence of a ssIAD continues to fall when greater numbers of pairs of readings are taken into account. Therefore, if a ssIAD is detected on the first measurement, we propose that BP should be taken simultaneously at least 3 times in both arms with the mean IAD calculated for the 2nd and 3rd readings in order to more accurately estimate a “true” IAD.

SUPPLEMENTARY MATERIAL

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

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REFERENCES

1. Weinberg I, Gona P, O’Donnell CJ, Jaff MR, Murabito JM. The systolic blood pressure difference between arms and cardiovascular disease in the Framingham Heart Study. Am J Med 2014; 127:209–215.
2. Clark CE, Taylor RS, Shore AC, Campbell JL. The difference in blood pressure readings between arms and survival: primary care cohort study. BMJ 2012; 344:e1327.
3. Clark CE, Taylor RS, Shore AC, Ukoumunne OC, Campbell JL. Association of a difference in systolic blood pressure between arms with vascular disease and mortality: a systematic review and meta-analysis. Lancet 2012; 379:905–914.
4. NICE. Hypertension: Clinical Management of Primary Hypertension in Adults London 2011. London: National Clinical Guideline Centre, The Royal College of Physicians.
5. Singh S, Sethi A, Singh M, Khosla S. Prevalence of simultaneously measured interarm systolic blood pressure difference and its clinical and demographic predictors: a systemic review and meta-analysis. Blood Press Monit 2015; 20:178–185.
6. Clark CE, Taylor RS, Shore A, Campbell J. Prevalence of systolic inter-arm differences in blood pressure for different primary care populations: systematic review and meta-analysis. Br J Gen Pract 2016; 66:e297–e308.
7. Abovyan V, Kamineni A, Allison MA, McDermott MM, Crouse JR, Ni H, Szklo M, Criqui MH. The epidemiology of subclavian stenosis and its association with markers of subclinical atherosclerosis: the Multi-Ethnic Study of Atherosclerosis (MESA). Atherosclerosis 2010; 211:266–270.
8. Kimura A, Hashimoto J, Watabe D, Takahashi H, Ohkubo T, Kikuya M, Imai Y. Patient characteristics and factors associated with inter-arm difference of blood pressure measurements in a general population in Osaka, Japan. J Hypertens 2004; 22:2277–2283.
9. Fotherby MD, Panayiotou B, Potter JF. Age-related differences in simultaneous interarm blood pressure measurements. Postgrad Med J 1993; 69:194–196.
10. Joshi P, Islam S, Pais P, Reddy S, Dorairaj P, Kazmi K, Pandey MR, Haque S, Mendis S, Rangarajan S, Yusuf S. Risk factors for early myocardial infarction in South Asians compared with individuals in other countries. JAMA 2007; 297:286–294.
11. Lip GY, Barnett AH, Bradbury A, Cappuccio FP, Gill PS, Hughes E, Imray C, Jolly K, Patel K. Ethnicity and cardiovascular disease prevention in the United Kingdom: a practical approach to management. J Hum Hypertens 2007; 21:183–211.
12. Anand SS, Yusuf S, Vukan V, Devanesen S, Teo KK, Montague PA, Kelemen L, Yi C, Lonn E, Gerstein H, Hegele RA, McQueen M. Differences in risk factors, atherosclerosis, and cardiovascular disease between ethnic groups in Canada: the Study of Health Assessment and Risk in Ethnic groups (SHARE). Lancet 2000; 356:279–284.
13. Mensah GA, Mokdad AH, Ford ES, Croft JB. State of disparities in cardiovascular health in the United States. Circulation 2005; 111:1233–1241.
14. Cappuccio FP, Cook DG, Atkinson RW, Strazzullo P. Prevalence, detection, and management of cardiovascular risk factors in different ethnic groups in south London. Heart 1997; 78:555–563.
15. Chiu M, Austin PC, Manuel DG, Tu JV. Comparison of cardiovascular risk profiles among ethnic groups using population health surveys between 1996 and 2007. CMAJ 2010; 182:E301–E310.
16. Howard VJ. Reasons underlying racial differences in stroke incidence and mortality. Stroke 2013; 44:S126–S128.
17. Sheng CS, Liu M, Zeng WF, Huang QF, Li Y, Wang JG. Four-limb blood pressure as predictors of mortality in elderly Chinese. Hypertension 2013; 61:1155–1160.
18. Kim KB, Oh MK, Kim HG, Ki JH, Lee SH, Kim SM. Inter-arm differences in simultaneous blood pressure measurements in ambulatory patients without cardiovascular diseases. Korean J Fam Med 2013; 34:98–106.
19. Verberk WJ, Kessels AG, Thiën T. Blood pressure measurement method and inter-arm differences: a meta-analysis. Am J Hypertens 2011; 24:1201–1208.
20. Lane D, Beeveres M, Barnes N, Bourne J, John A, Malins S, Beeveres DG. Inter-arm differences in blood pressure: when are they clinically significant? J Hypertens 2002; 20:1089–1095.
21. Wood S, Martin U, Gill P, Greenfield SM, Haque MS, Mant J, Mohammed MA, Heer G, Johal A, Kaur R, Schwartz C, McNamur RJ. Blood pressure in different ethnic groups (BP-Eth): a mixed methods study. BMJ Open 2012;2:e001598.
22. McKenzie K, Crowcroft N. Ethnicity, race, and culture: guidelines for research, audit, and publication. BMJ 1996; 312:1094.
23. Wright JM, Mattu GS, Perry Jr TL, Gelferc ME, Zorn A, McKenzie K, Crowcroft N. Ethnicity, race, and culture: guidelines for research, audit, and publication. BMJ 1996; 312:1094.
24. Baumgart P, Kamp J. Accuracy of the SpaceLabs Medical 90217 ambulatory blood pressure monitor. Blood Press Monit 2001; 6:161–165.
25. Rasmussen SL, Torp-Pedersen C, Borch-Johnsen K, Ibsen H. Normal white-coat hypertension response: prevalence and predictors. Blood Press Monit 2015; 20:178–185.
26. Verberk WJ, Kessels AG, Thiën T. Blood pressure measurement method and inter-arm differences: a meta-analysis. Am J Hypertens 2011; 24:1201–1208.
27. NRS. Blood pressure in different ethnic groups (BP-Eth): a mixed methods study. BMJ Open 2012;2:e001598.
28. McKenzie K, Crowcroft N. Ethnicity, race, and culture: guidelines for research, audit, and publication. BMJ 1996; 312:1094.
29. Wright JM, Mattu GS, Perry Jr TL, Gelferc ME, Zorn A, McKenzie K, Crowcroft N. Ethnicity, race, and culture: guidelines for research, audit, and publication. BMJ 1996; 312:1094.
30. Altman DG. Practical Statistics for Medical Research. Chapman & Hall: London, 1991.
31. Joshi S, Christopher C, Campbell J. What is the normal inter-arm difference? It depends on left or right handedness: systematic review and meta-analysis. J Hypertens 2016/17; 34(10):1421–1427.
32. Rasmussen SL, Torp-Pedersen C, Borch-Johnsen K, Ibsen H. Normal white-coat hypertension response: prevalence and predictors. Blood Press Monit 2001; 6:161–165.
33. NHS England. 2016/17 General Medical Services (GMS) contract Quality and Outcomes Framework (QOF) Guidance for GMS contract 2016/17. London: NHS Employers, 2016.
34. Martin U, Holder R, Hodgkinson J, McNamur RJ. Inter-arm blood pressure differences compared with ambulatory monitoring: a manifestation of the ‘white-coat’ effect? Br J Gen Pract 2013; 63:e97–103.
31. van der Hoeven NV, Lodestijn S, Nanninga S, van Montfrans GA, van den Born BJ. Simultaneous compared with sequential blood pressure measurement results in smaller inter-arm blood pressure differences. J Clin Hypertens (Greenwich) 2013; 15:839–844.

32. NHS England. Health Survey for England—Health, Social Care and Lifestyles 2011 <http://digital.nhs.uk/catalogue/PUB09300/HSE2011-Ch3-Hypertension.pdf> 2016. Access 26 September 2016.

33. Mancia G, Fagard R, Narkiewicz K, Redon J, Zanchetti A, Böhm M, Christiaens T, Cifkova R, De Backer G, Dominiczak A, Galetska M, Grobbee DE, Jaarsma T, Kirchhof P, Kjeldsen SE, Laurent S, Manolis AJ, Nilsson PM, Ruilope LM, Schmidt RE, Sirnes PA, Sleight P, Viigimaa M, Waerbel A, Zannad F, Redon J, Dominiczak A, Narkiewicz K, Nilsson PM, Burnier M, Viigimaa M, Ambrosioni E, Cauterfield M, Coca A, Olsen MH, Schmidt JE, Tsioftis C, van de Borne P, Zamorano JL, Achenbach S, Baumgartner H, Bax JJ, Bueno H, Dean V, Deaton C, Erol C, Fagard R, Ferrari RI, Haselbauer D, Hoes AW, Kirchhof P, Knutsinis J, Kolb P, Lancellotti P, Linhart A, Nihoyannopoulos P, Piepoli MF, Ponikowski P, Sirnes PA, Tamargo JL, Tendera M, Torbicki A, Wijn W, Windecker S, Clement DL, Coca A, Gillebert TC, Tenden M, Rosie EA, Ambrosioni E, Anker SD, Bauersachs J, Hitzh JB, Caulifield M, De Buyzere M, De Geest S, Derumaux GA, Erdine S, Farsang C, Funck-Brentano C, Gerc V, Germino G, Gielen S, Haller H, Hoes AW, Jordan J, Kan T, Komajda M, Lovic M, Mahrholdt H, Olsen MH, Ostergren J, Parati G, Perk J, Polsonia J, Pospelova BA, Reiner Z, Ryden L, Sirekno Y, Stanton A, Struijker-Boudier H, Tsiouris C, van de Borne P, Winodecuolas C, Volpe M, Wood DA. 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). Eur Heart J 2013; 34:2159–2219.

34. English JA, Carell ES, Guidera SA, Tripp HF. Angiographic prevalence and clinical predictors of left subclavian stenosis in patients undergoing diagnostic cardiac catheterization. Catheter Cardiovasc Interv 2001; 54:8–11.

35. Sun H, Li P, Su H, Wang J, Hu W, Li J, Liu Y, Cheng X. The detection rates of inter-arm systolic blood pressure difference vary with blood pressure levels in hypertensive patients under antihypertensive therapy. Int J Cardiol 2014; 172:e419–e420.

36. Head GA, Mihalidou AS, Duggan KA, Belin LJ, Berry N, Brown MA, Bune AJ, Cowley D, Chalmers JP, Howe PR, Hodgson J, Ludbrook J, Mangoni AA, McGrath BP, Nelson MR, Sharman JE, Stowasser M; Ambulatory Blood Pressure Monitoring Group of the High Blood Pressure Research Council of Australia. Definition of ambulatory blood pressure targets for diagnosis and treatment of hypertension in relation to clinic blood pressure: prospective cohort study. BMJ 2010; 340:c1104.

37. Lohmann FW, Eckert S, Verbek WR. Interarm differences in blood pressure should be determined by measuring both arms simultaneously with an automatic oscillometric device. Blood Press Monit 2011; 16:37–42.

38. Eguchi K, Yacoub M, Jalal J, Gerin W, Schwartz JE, Pickering TG. Consistency of blood pressure differences between the left and right arms. Arch Intern Med 2007; 167:388–393.

39. Staessen JA, O’Brien E, Amery AK, Atkins N, Baumgart P, De Cort P, Degaute JP, Dolenc P, De Gaudemas R, Enstrom I. Ambulatory blood pressure in normotensive and hypertensive subjects: results from an international database. J Hypertens Suppl 1994;12:S13–S22.

40. Thomas O, Shipman KE, Day K, Thomas M, Martin U, Dasgupta I. Prevalence and determinants of white coat effect in a large UK hypertension clinic population. J Hum Hypertens 2016; 30:386–391.