Sleep Duration and Ambulatory Blood Pressure in Black and White Adolescents

Elizabeth J. Mezick, Martica Hall, Karen A. Matthews

Abstract—Self-reported short sleep duration is linked to higher blood pressure and incident hypertension in adults. Few studies have examined sleep and blood pressure in younger samples. We evaluated the associations between actigraphy-assessed time spent asleep and ambulatory blood pressure in adolescents. Participants were 246 black and white adolescents (mean age: 15.7 years) who were free from cardiovascular or kidney disease and were not taking sleep, cardiovascular, or psychiatric medications. Sleep duration and efficiency were assessed with in-home wrist actigraphy and sleep diaries across 1 week; ambulatory blood pressure monitoring was used to obtain 24-hour, sleep, and wake blood pressure, as well as sleep:wake blood pressure ratios across 2 full days and nights. Results showed that shorter actigraphy-assessed sleep across 1 week was related to higher 48-hour blood pressure and higher nighttime blood pressure. Shorter sleep was also related to a higher systolic blood pressure sleep:wake ratio. These results were independent of age, race, sex, and body mass index. Follow-up analyses by race revealed that associations between sleep duration and blood pressure were largely present in white, but not black, adolescents. These data are consistent with the hypothesis that the cardiovascular consequences of short sleep may begin as early as adolescence. (Hypertension. 2012;59:747-752.)

Key Words: ambulatory blood pressure ■ sleep duration ■ actigraphy ■ adolescent ■ race

Reports of usual sleep duration are inversely linked to resting blood pressure (BP) and hypertension risk in adults.1–4 For instance, in the First National Health and Nutrition Examination Survey, persons reporting usual sleep duration of ≤5 hours per night were at increased risk of hypertension over the 8- to 10-year follow-up period, after adjusting for a variety of sociodemographic and health factors.5 Correlations between self-reports of typical sleep duration and behavioral measures of sleep are modest.5 Thus, actigraphy that infers sleep duration based on nocturnal movement is a useful complement to subjective reports. Of the 2 studies that have examined actigraphy-assessed sleep and BP in adults, 1 reported that short sleep was related to higher BP and increased hypertensive risk in middle-aged adults,6 whereas the other failed to detect an association between sleep duration and hypertension in the elderly.7 Few studies have examined whether the link between sleep and BP is present in childhood or adolescence. BP levels in early life track into adulthood8 and may predict metabolic syndrome and carotid intima-media thickness decades later.9–13 Developmental changes in a variety of biological, social, and environmental factors in adolescence result in a large number of young people being exposed to a continual state of insufficient sleep.14 Thus, adolescence provides a critical period for studying the physiological consequences of chronic sleep deprivation, as well as a potential opportunity for prevention or early intervention for hypertension risk. The 1 study of actigraphy and BP in adolescents measured sleep across ≥3 weeknights and classified participants as prehypertensive based on an average of clinic BP measurements on a single day.15 In that study, low sleep efficiency (percentage of time in bed spent asleep) was associated with a 3.5-fold increase in the odds of prehypertension after adjustment for body mass index (BMI) and other covariates.15 Relationships of short sleep duration with prehypertension and systolic BP (SBP) were reduced to statistical trends after adjustment for BMI and other covariates. With regard to younger samples, no association was observed between actigraphy-measured sleep over 6 nights and 24-hour ambulatory BP levels in Finnish 8-year olds,16 and the link between short sleep, as assessed by waist actigraphy for 24 hours, and higher resting BP in New Zealand 7-year olds was attenuated after adjustment for BMI.17

In this study, we examined actigraphy measures of sleep duration and efficiency across 1 week in relation to 48 hours of ambulatory BP in a sample of black and white adolescents. There are several benefits of examining ambulatory BP relative to clinic BP, including increased reliability and a stronger approximation of cardiovascular risk.18 Use of ambulatory monitoring also permits measures of circadian BP pattern via separate day and night estimates, as well as the sleep:wake BP ratio, which predicts cardiovascular outcomes independent of daytime BP.19–21 In adult samples, higher sleep:wake BP ratios, or nondipping, are associated with sleep fragmentation and sleep disturbance.22–24 Our second-
ary aim was to investigate whether the links between sleep duration or efficiency and BP are similar in black and white adolescents, given that blacks are at increased risk for hypertension and have higher sleep:wake BP ratios and shorter sleep than whites in adult samples.25,26 We hypothesized that short, inefficient sleep would be related to higher 24-hour ambulatory BP and higher sleep:wake BP ratios, and we examined whether these associations varied by race.

Methods

Participants

We recruited 250 adolescents between the ages of 14 and 19 years from a local high school within 15 miles of downtown Pittsburgh. The high school served a diverse population, both in terms of ethnicity and socioeconomic status. Approval of the research project was obtained from the University of Pittsburgh Institutional Review Board. Participants, and in the case of students under the age of 18 years, a parent or legal guardian provided written informed consent before any research procedures. Exclusionary criteria included cardiovascular or kidney disease, medication use for emotional or psychological disorders, use of diabetes mellitus or BP medication, use of any medication known to affect the cardiovascular system or sleep, and inability to obtain BP readings (usually because of extreme obesity). Sixteen students who were screened were ineligible to participate because of taking medication that could affect study variables, and 7 students who signed consent did not begin the study.

Procedure

The study protocol lasted 1 week. Participants wore the ambulatory monitor for the 2 school days and nights and wore an Actigraph on their nondominant wrist continuously over the 7 days and nights of the study. They answered several questions about their sleep in a handheld computer each morning and evening. Additional study procedures included assessment of anthropometric measures, fasting venous blood draw, and psychosocial questionnaires and interview. After completion of the protocol, participants were compensated $100, and a follow-up report of the student’s BP, sleep, anthropometric, and lipid levels was sent to the student and his or her parent or guardian.

Measures

Ambulatory BP

Ambulatory BP was measured using the SpaceLabs model 90217 ambulatory BP monitor (Issaquah, WA), with pediatric, adult, or thigh cuff used depending on body size. The BP monitors were worn for 2 school days and nights and were programmed to take BP readings every 30 minutes during the day (from 7:00 AM to 10:00 PM) and hourly during the night (from 10:00 PM to 7:00 AM). Participants were told to wear the monitor at all times except when bathing. BPs that fell within the participant’s self-reported sleep window were used to calculate average nighttime BP based on a minimum of 5 readings across the night. BP readings that were ≥3 SDs from a participant’s individual mean were excluded. Outcome variables were as follows: (1) 24-hour BP, defined as the average of all of the valid BP readings obtained across the 2- to 3-day period; (2) nighttime BP, defined as the average of all of the valid BP readings obtained within the nighttime hours that the participant reported that he or she was attempting to sleep; (3) daytime BP, defined as the average of all of the valid BP readings during reported waking hours; and (4) the sleep: wake BP ratio, defined as the ratio of average nighttime BP to average daytime BP. To examine effects of potential clinical relevance, we also included 24-hour prehypertension, defined as an average SBP >120 mm Hg or diastolic BP (DBP) >80 mm Hg over the entire study (n = 109).27 Using the recommended adult cutoff of daytime SBP >135 mm Hg or DBP >85 mm Hg captured a small subset of the sample (n = 20),18 so we reduced the cutoff for elevated daytime BP to include those with average daytime SBP >130 mm Hg or DBP >80 mm Hg (N = 57). BP nondipping was defined as <10% decline in sleep SBP relative to wake SBP or <10% decline in sleep DBP relative to wake DBP (n = 68).21

Sleep

The Mini-mitter actiwatch model AW-16 (Phillips Respironics, Bend, OR) collected actigraphy data continuously over 7 days and nights. Actigraphs were configured to collect data over a 1-minute epoch. Stored data were downloaded into the Actiware software program (version 5.57) for processing and analysis. The medium threshold (default) was selected to detect nocturnal sleep periods of ≥3 hours in duration based on sleep onset and offset using the 10-minute criteria. Each night’s data were reviewed individually and revealed that many sleep periods occurred before or after the recorded diary bedtimes and awake times. Thus, sleep periods occurring within 30 minutes of the major nocturnal sleep interval (either 30 minutes before sleeping or after waking) that were ≥15 minutes in duration were combined with the major sleep interval (ie, if a 6-hour sleep interval was detected from 12:00 AM to 6:00 AM, and a 20-minute sleep interval was detected beginning at 11:30 PM, the 20-minute interval was combined with the major sleep interval; the new major rest interval would become 11:30 PM to 6:00 AM). All of the subsequent sleep variables were then calculated from data within these set rest periods. Total sleep time was calculated as the time spent asleep between initial sleep onset and final sleep offset, excluding periods of wakefulness throughout the night. Sleep efficiency was calculated as the percentage of time attempting to sleep that was actually spent asleep. The actiwatch has been widely used in research studies and has been validated against polysomnography measures in clinic.28,29

Diary total sleep time was calculated as the difference between the time that one reported trying to fall asleep and the time that he or she reported finally waking, minus the self-reported number of minutes it took to fall asleep and the number of minutes spent awake after sleep onset. All of the sleep measures were averaged across the 7 nights of the study.

Covariates

Age, sex, and race/ethnicity were determined by self-report. Height was measured using a stadiometer, and weight was measured on a Tanita digital scale to the tenth of a pound. BMI was calculated using the National Heart, Lung, and Blood Institute online calculator.

Analysis

Data were checked for normality. Two participants had BMI values that fell >4 SDs from the mean and were excluded from analyses. Two participants were excluded because of equipment failure or misuse. Therefore, analyses examining daytime and 24-hour ambulatory BPs were based on a total of 246 participants. An additional 2 participants had incomplete nocturnal BP data because of taking off the cuff on ≥1 night; therefore, analyses examining nocturnal BP or sleep: wake BP ratios were based on a total of 244 participants. We examined the associations by linear regression between total sleep time and sleep efficiency with continuous ambulatory BP outcomes, including 24-hour BP, daytime BP, nighttime BP, and sleep: wake BP ratios. Standardized βs are reported, with coefficients signifying the SD change in BP as a result of a 1-SD increase in sleep duration in hours (1 SD = 0.79 hours or 47.4 minutes). Associations with categorical cutoffs of elevated BP were tested in logistic regression models in 1-hour units of sleep. Finally, we included a race-total sleep time interaction term in all of the above models to test for differences in the links between sleep duration and efficiency with BP between white and black adolescents. All of the base models were adjusted for age, sex, and race. BMI was added in the final step to examine whether it attenuated associations between sleep and BP. Results were also adjusted for BMI percentile and revealed identical results.

Results

Sample characteristics are shown in Table 1. Slightly more than half of the sample were female (53.3%), and slightly
more than half were black (56.5%) (Table 1). Average time spent asleep across the 7 nights of the study by actigraphy was 6.45 hours (SD 0.79; range: 4.27–9.22), and average sleep efficiency was 82.66% (SD 4.87; range: 67.78–94.11). On average, ambulatory nighttime BP declined by 13% for SBP and by 20% for DBP. After adjustment for age, sex, and BMI, black adolescents had higher 24-hour DBP, nighttime SBP, and sleep: wake SBP ratios.  

**Actigraphy-Measured Sleep and BP**

Shorter sleep duration across the study was associated with higher 24-hour SBP and DBP across the 2 days of ambulatory monitoring (Table 2; Figure). The relationship between sleep and 24-hour BP levels was driven largely by nocturnal BP; shorter sleep was related to higher nighttime SBP and DBP. Sleep duration tended to be associated with daytime BP in the hypothesized direction, but these relationships were not statistically significant. Finally, there was an inverse association between sleep duration and sleep:wake SBP ratios, such that shorter sleep was related to a smaller nocturnal decline in BP. All of the associations between sleep duration and BP remained after adjusting for BMI. Sleep duration was not related to sleep:wake DBP ratios.

**Table 1. Sample Characteristics by Race**

<table>
<thead>
<tr>
<th>Variable</th>
<th>Black (n=139)</th>
<th>White (n=107)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No. (%) Mean (SD) Range</td>
<td>No. (%) Mean (SD) Range</td>
</tr>
<tr>
<td>Male</td>
<td>49 (45.8) 15.7 (1.3) 14–19</td>
<td>66 (47.5) 15.7 (1.3) 14–19</td>
</tr>
<tr>
<td>Age, y</td>
<td>15.7 (1.3) 14–19</td>
<td>15.7 (1.3) 14–19</td>
</tr>
<tr>
<td>Body mass index</td>
<td>26.1 (5.5) 17.8–43.5</td>
<td>26.2 (6.5) 16.4–46.9</td>
</tr>
<tr>
<td>Body mass index percentile</td>
<td>80.7 (19.5) 10–100</td>
<td>76.3 (25.9) 3–100</td>
</tr>
<tr>
<td>Total sleep time, h</td>
<td>6.2 (0.69)* 4.3–7.7</td>
<td>6.7 (0.83)* 4.9–9.2</td>
</tr>
<tr>
<td>≥5 h total sleep time</td>
<td>6 (4.3)</td>
<td>1 (0.9)</td>
</tr>
<tr>
<td>≥6 h total sleep time</td>
<td>43 (30.9) 17 (15.9)</td>
<td>53 (49.5)</td>
</tr>
<tr>
<td>≥7 h total sleep time</td>
<td>70 (50.4)</td>
<td>53 (49.5)</td>
</tr>
<tr>
<td>≥8 h total sleep time</td>
<td>20 (14.4) 28 (26.2)</td>
<td></td>
</tr>
<tr>
<td>≤10 h total sleep time</td>
<td>0 (0) 8 (7.5)</td>
<td></td>
</tr>
<tr>
<td>24-h ambulatory SBP</td>
<td>119.7 (8.7)‡ 97.0–144.3</td>
<td>117.7 (9.0)‡ 97.3–148.9</td>
</tr>
<tr>
<td>24-h ambulatory DBP</td>
<td>69.7 (6.0)† 54.2–82.2</td>
<td>68.3 (5.3)† 58.0–79.1</td>
</tr>
<tr>
<td>Daytime SBP</td>
<td>122.6 (9.2)† 93.5–149.5</td>
<td>121.0 (9.1)† 101.7–154.2</td>
</tr>
<tr>
<td>Daytime DBP</td>
<td>72.6 (6.3) 53.8–86.2</td>
<td>71.4 (5.6) 58.9–82.8</td>
</tr>
<tr>
<td>Nighttime SBP</td>
<td>107.1 (8.3)† 87.0–137.2</td>
<td>104.2 (9.3)† 86.2–128.8</td>
</tr>
<tr>
<td>Nighttime DBP</td>
<td>57.4 (5.6)‡ 43.6–72.3</td>
<td>56.3 (5.4)‡ 45.2–70.6</td>
</tr>
<tr>
<td>Sleep:wake SBP Ratio</td>
<td>0.88 (0.06)† 0.75–1.10</td>
<td>0.86 (0.05)† 0.70–0.99</td>
</tr>
<tr>
<td>Sleep:wake DBP Ratio</td>
<td>0.79 (0.07) 0.59–1.04</td>
<td>0.79 (0.06) 0.59–0.98</td>
</tr>
</tbody>
</table>

24-h prehypertension (24-h SBP ≥120 mm Hg or DBP ≥80 mm Hg) 65 (46.8) 44 (41.1)  
Elevated daytime BP (day SBP ≥130 mm Hg or DBP ≥80 mm Hg) 34 (24.5) 23 (21.5)  
Nondipping (SBP or DBP sleep:wake ratio ≥0.90) 46 (33.3)† 22 (20.8)†  

BP indicates blood pressure; SBP, systolic blood pressure; DBP, diastolic blood pressure.  
*P<0.01, with differences by race after adjusting for age, sex, and body mass index.  
†P<0.05, with differences by race after adjusting for age, sex, and body mass index.  
‡P<0.10, with differences by race after adjusting for age, sex, and body mass index.

**Table 2. Results From Linear Regression Models of Total Sleep Time and Ambulatory BP**

<table>
<thead>
<tr>
<th>Variable</th>
<th>24-Hour</th>
<th>Daytime</th>
<th>Nighttime</th>
<th>Sleep/Wake Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SBP</td>
<td>DBP</td>
<td>SBP</td>
<td>DBP</td>
</tr>
<tr>
<td>Total sleep time</td>
<td>β</td>
<td>P</td>
<td>β</td>
<td>P</td>
</tr>
<tr>
<td></td>
<td>−0.17</td>
<td>0.01</td>
<td>−0.11</td>
<td>0.09</td>
</tr>
<tr>
<td>Race×total sleep time</td>
<td>0.97</td>
<td>0.05</td>
<td>0.82</td>
<td>0.10</td>
</tr>
<tr>
<td>Total sleep time: blacks</td>
<td>−0.02</td>
<td>0.80</td>
<td>−0.04</td>
<td>0.60</td>
</tr>
<tr>
<td>Total sleep time: whites</td>
<td>−0.34</td>
<td>0.01</td>
<td>−0.32</td>
<td>0.02</td>
</tr>
</tbody>
</table>

Covariates for full sample analyses include age, sex, race, and body mass index; covariates for race-stratified analyses include age, sex, and body mass index. Standardized β-coefficients signify the SD change in blood pressure as a result of a 1-SD increase in total sleep time (1 SD = 0.79 h or 47.4 min). SBP indicates systolic blood pressure; DBP, diastolic blood pressure.
Adolescents sleeping for a shorter duration were more likely to be classified as prehypertensive based on 24-hour BP (average 24-hour SBP >120 mm Hg or DBP >80 mm Hg; Table 3). In a separate analysis using the cutoff of daytime SBP >130 mm Hg or DBP >80 mm Hg, shorter sleepers were also more likely to be classified as having elevated daytime BP. Using the criterion of <10% decline in nocturnal SBP or DBP relative to daytime levels, adolescents with shorter sleep were more likely to be categorized as nondippers. All of the associations between sleep duration and BP cutoffs remained after adjusting for BMI. Sleep also showed a smaller decline in nocturnal SBP relative to daytime. Short sleep was associated with increased odds of meeting prehypertensive levels across a 24-hour period and average daytime SBP >130 mm Hg or DBP >80 mm Hg across 2 school days. Shorter sleepers were also more likely to be classified as nondippers relative to those getting more sleep. Although previous data have suggested a potential mediating role of BMI in the link between sleep and BP, all of the associations in the current sample were independent of BMI.

Actigraphy-assessed sleep efficiency, an index of sleep continuity, was unrelated to ambulatory BP in this sample. This is in contrast to the finding of Javaheri et al that sleep efficiency, rather than sleep duration, was more consistently linked with in-clinic BP in the Cleveland Children’s Sleep and Health Study. One reason for this discrepancy may be sample differences in sleep characteristics. Mean sleep duration and efficiency in the current study were relatively low compared with the Cleveland sample, which included a similar racial/ethnic composition but was slightly younger. Although only 25% of adolescents in the Cleveland study were classified as having a sleep efficiency ≤85%, more than two thirds of adolescents in the current sample met that criterion. Among adults in the Coronary Artery Risk Development in Young Adults Study, both actigraphy-assessed short sleep duration and low sleep maintenance (an index similar to sleep efficiency) were associated with increased BP; however, only short sleep duration predicted increased risk of incident hypertension.

Our results were by and large present in white, but not black, adolescents. It is possible that a restricted range of sleep duration among black adolescents inhibited our ability to detect an association with BP (only 14% exceeded 7 hours of sleep measured by actigraphy across the week compared with 30% of whites). The Cleveland Sleep and Health Study

### Table 3. Results From Logistic Regression Models of Total Sleep Time and Ambulatory BP

<table>
<thead>
<tr>
<th>Variable</th>
<th>24-h Prehypertension (SBP &gt;120 mm Hg or DBP &gt;80 mm Hg; n=109)</th>
<th>Elevated Daytime BP (SBP &gt;130 mm Hg or DBP &gt;80 mm Hg; n=57)</th>
<th>BP Nondipping (SBP or DBP sleep:wake ratio &gt;0.90; n=68)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total sleep time</td>
<td>OR 0.66 95% CI 0.46–0.97</td>
<td>OR 0.65 95% CI 0.42–0.98</td>
<td>OR 0.66 95% CI 0.44–0.99</td>
</tr>
<tr>
<td>Total sleep time: blacks</td>
<td>OR 1.04 95% CI 0.63–1.73</td>
<td>OR 1.03 95% CI 0.59–1.82</td>
<td>OR 0.83 95% CI 0.48–1.45</td>
</tr>
<tr>
<td>Total sleep time: whites</td>
<td>OR 0.36 95% CI 0.19–0.68</td>
<td>OR 0.27 95% CI 0.12–0.61</td>
<td>OR 0.56 95% CI 0.29–1.06</td>
</tr>
</tbody>
</table>

OR indicates odds ratio for meeting the outcome criterion, given a 1-h increase in total sleep time; BP, blood pressure; SBP, systolic blood pressure; DBP, diastolic blood pressure. Covariates for full-sample analyses include age, sex, race, and body mass index; covariates for race-stratified analyses include age, sex, and body mass index.
did not test for race differences in their findings, and, in Coronary Artery Risk Development in Young Adults Study, race did not modify associations between sleep duration or sleep maintenance and BP. With regard to other cardiovascular risk factors, some studies have reported that short sleep is more closely linked to inflammatory biomarkers in black adults than in whites, whereas others have failed to detect a difference in links between sleep duration and traditional cardiovascular risk factors or coronary calcification by race. Although the interaction analyses in this sample were exploratory and did not reach statistical significance in all of the cases, the overall pattern of effects suggests that ethnic variation in sleep and BP may warrant further study.

Several physiological mechanisms could link shorter sleep to higher BP. Acute total sleep deprivation studies in healthy young adults produce an increase in DBP along with decreased muscle sympathetic nerve activity, consistent with a shift in the arterial baroreflex toward a higher BP. Experimental sleep loss also can cause increases in vascular endothelial adhesion markers, followed by elevated BP. Partial sleep deprivation over several nights has been linked to changes in sympathovagal activity, as evidenced by decreased heart rate variability and increased catecholamine release. Thus, small amounts of sleep loss accrued over time could potentially result in similar pathophysiological effects, leading to increased BP. The association between sleep duration and 24-hour BP was largely attributed to higher nocturnal BP. Previous work has shown that parasympathetic activity increases across the night, reaching a peak in the fourth nonrapid eye movement sleep period. It is possible that those with curtailed sleep do not reach this sleep stage and the concurrent increase in parasympathetic activity and, thus, exhibit increased nocturnal pressure. An additional possibility is that higher nocturnal BP and nondipping reflect elevated nocturnal activity among shorter sleepers. This is less likely given that nocturnal BP measurements were based on reported sleep hours, which capture individuals’ sleep more accurately than an arbitrary definition of nighttime and sleep efficiency was unrelated to nighttime BP. However, variation in sleep architecture not captured by actigraphy, such as less time spent in deep sleep, could contribute to higher nocturnal BP.

A limitation of this study was our inability to assess sleep-disordered breathing. Obstructive sleep apnea is associated with increased daytime and nighttime ambulatory BP in children and adolescents in a number of studies, although the overall literature is mixed. and actigraphy may misestimate total sleep time among those with sleep-disordered breathing. Thus, sleep apnea may be a confounder of the observed relationships. The cross-sectional design precludes implications about causation, and it is possible that underlying autonomic dysfunction resulted in both shorter sleep and elevated BP. A substantial number of participants had ambulatory BP during the day at prehypertensive levels, in part because of the high BMI of the participants. Thus, the findings would not generalize to samples with fewer obese individuals. Finally, the reasons for individual amounts of sleep duration are unknown, and voluntary (ie, time demands) versus involuntary (ie, inability to fall asleep) sleep curtail-

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