Alcohol Consumption and Vascular Variability Anomalies (VVAs) and Disorders (VVDs) Detected by C-ABPM

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Abstract: Alterations of the circadian rhythms in blood pressure (BP) and heart rate (HR) have been associated with an increased cardiovascular disease risk. The influence of alcohol consumption on the circadian rhythm characteristics of BP is here examined, and the extent to which such alteration of the circadian BP rhythm contributes to cardiovascular disease risk is assessed.

Keywords: Alcohol Consumption, Vascular Variability Anomalies (VVAs), C-ABPM, Relative Risk.

INTRODUCTION

Blood pressure (BP) and heart rate (HR) are highly variable. Their variability includes partly built-in rhythms. Circadian rhythms in particular are generally prominent [1], usually with values lower during sleep/rest, a small increase around mid-sleep [2] followed by a larger increase after awakening, a post-prandial dip accentuating with age [3], and a slow decrease in the evening. A host of other external factors also influence BP and HR, including emotions [4], exercise [5] and other lifestyle features such as smoking [6].

Meals have also been shown to affect BP and HR. Indeed, in a study of 164 newborns fed every 3 hours (except during one timepoint at night) and monitored automatically around the clock during the first week of life, an about 3-hour component prominently characterized HR [7]. In another study where babies were fed every 4 hours, a statistically significant spectral peak was found at a frequency of 1 cycle in 4 hours [7]. An increase in HR following meals was demonstrated in a study of clinically healthy nurses, 18-25 years of age, in Tokyo, Japan [8]. They were monitored at 15-minute intervals for 48 hours, one group of 44 nurses in November 1987 and another group of 50 nurses in June 1988. Time-specific reference limits (chronodesms) computed as 90% prediction limits showed good reproducibility, notably in terms of the presence of 3 peaks in the morning, around noon, and in the evening, coinciding with meal times [8]. Such meal-associated peaks were not found in similar chronodesms computed from other populations of healthy individuals who did not follow a synchronized a living routine as the Japanese nurses. Figs. (1-3) visualize the circadian patterns as means +/- 1 standard error for the two groups, documenting the reproducibility of the 3 daytime peaks with the nighttime drop in BP and HR.

Outcome studies [9-14] have shown that cardiovascular disease risk is increased not only in association with a high BP, but also with alterations of the variability in BP and/or HR [9, 10, 15, 16]. In particular, an excessive circadian amplitude of BP (above the upper 95% prediction limit of clinically healthy peers matched by gender and age), a condition called CHAT (brief for Circadian Hyper-Amplitude-Tension) was associated with a large increase in the risk, larger that that of a high BP itself, Fig. (4). Contrary to the linear relation of risk with respect to the BP MESOR (Midline Estimating Statistic Of Rhythm, a rhythm-adjusted mean), the relation of cardiovascular disease risk to the circadian amplitude of BP is nonlinear; risk is increased only after the circadian amplitude of BP exceeds a threshold, Fig. (5).

Herein, we examine whether alcohol affects the circadian rhythm of BP and assess any associated increase in cardiovascular disease risk.

SUBJECTS AND METHODS

The BP and HR of 297 Japanese non-diabetic patients (145 women and 152 men) were automatically measured every 15 minutes for 48 hours with an ambulatory monitor at the start of the study [9, 10]. There were 121 MESOR-normotensive subjects and 176 treated MESOR-hypertensive patients. Outcomes (cerebral ischemic event, coronary artery disease, nephropathy, and retinopathy), absent at the start of the study, were checked every 6 months for 6 years.

Each record was analyzed chronobiologically [17, 18]. Estimates of the MESOR and of the circadian amplitude and acrophase were interpreted in the light of reference values from clinically healthy peers matched by gender and age. Screening for VVDs identified patients with MESOR-Hypertension (MH) and CHAT, among others. The relative risk [19] associated with MH and CHAT was determined and compared with that of other known risk factors assessed concomitantly. These included obesity, high cholesterol, gender, family history, smoking, age, as well as alcohol consumption. The relative risk of CHAT was further
assessed in patients who checked negative for any one of the other risk factors considered each separately.

Previously, relative risk had been assessed separately for each kind of outcome [9, 10, 20]. Notably in the case of cerebral ischemic events and nephropathy, CHAT was found to represent, at least numerically, a risk higher than that of MESOR-Hypertension. As compared to obesity, high cholesterol, male gender, a positive family history (FH) of high BP and/or related cardiovascular disease, smoking, being older than 60 years of age, and MESOR-Hypertension, CHAT carried a higher risk of cerebral ischemic events and nephropathy. When considering subpopulations not presenting with one of the other given risk factors, the risk associated with CHAT remained statistically significant, indicating that the risk of cerebral ischemic events and nephropathy was raised in the presence of CHAT, irrespective of the effect of the other risk factor being tested concomitantly [20].
Herein, relative risk is assessed in relation to all outcomes considered jointly. After 6 years, 39 patients developed coronary artery disease, cerebral ischemia, nephropathy and/or retinopathy.
RESULTS

Among the 297 patients, 195 were drinkers and 102 non-drinkers. Adverse outcomes after 6 years affected 18.6% of drinkers as compared to 10.3% of non-drinkers ($\chi^2=4.114$, $P=0.043$). The relative risk (RR) of alcohol consumption with its 95% confidence interval (CI) is $1.82 \ [1.02, 3.25]$. As seen in Fig. (6), alcohol consumption is associated with a statistically significant increase in cardiovascular disease risk, whereas in the population investigated, obesity, high cholesterol, male gender, and a positive family history of high BP and/or related cardiovascular disease are not. Smoking is associated with an even higher RR of 2.33 $[1.31, 4.13]$ ($\chi^2=8.425$, $P=0.004$), as is age above 60 years with a RR of 3.64 $[2.08, 6.36]$ ($\chi^2=21.145$, $P<0.001$), and MH with a RR of 8.25 $[2.60, 26.18]$ ($\chi^2=20.310$, $P<0.001$). Systolic and diastolic CHAT also have a relative risk statistically significantly higher than 1 ($P<0.001$). Whereas it is less than that of MH, their smaller CIs indicate a higher specificity [21].

Similar results are observed for the subpopulation of non-drinkers, as shown in Fig. (7). No statistically significant increase in cardiovascular disease risk is found in association with obesity, high cholesterol, male gender or a positive FH. Smoking has a RR of 2.27 $[0.90, 5.69]$ reaching borderline statistical significance ($P=0.085$). Age (RR=6.63, $P<0.001$) and MH (RR=7.56, $P=0.001$) remain the largest risk factors, while diastolic CHAT remains a significant risk factor (RR=3.23, $P=0.019$), systolic CHAT reaching borderline statistical significance (RR=2.62, $P=0.059$). In the smaller subpopulation of drinkers, only MH and CHAT are found to significantly increase cardiovascular disease risk (MH: RR=8.23, $P=0.007$; S-CHAT: RR=3.38, $P=0.003$; D-CHAT: RR=4.83, $P<0.001$), Fig. (8). Again, the lesser RR of CHAT
**Fig. (6).** Overall cardiovascular disease risk (diamonds) is statistically significantly increased in association with alcohol consumption, as evidenced by the non-overlap of equal risk (RR=1) by the 95% confidence interval of the relative risk (horizontal bars). Apart from age, MESOR-Hypertension, and CHAT, only smoking is also associated with a statistically significant increase in cardiovascular disease risk in the clinic population of 297 Japanese patients. © Halberg (with permission).

**Fig. (7).** Among the 195 non-drinkers, age, MESOR-Hypertension and CHAT remain statistically significantly associated with an increase in overall cardiovascular disease risk, while smoking reaches borderline statistical significance. © Halberg (with permission).
Fig. (8). In the smaller subpopulation of 102 drinkers, only MESOR-Hypertension and CHAT are associated with a statistically significant increase in overall cardiovascular disease risk. © Halberg (with permission).

Fig. (9). Alcohol consumption is associated with a small statistically significant increase in the BP MESOR, with a much larger statistically significant increase in the circadian amplitude of BP, but no effect on pulse pressure. © Halberg (with permission).

as compared to MH may be viewed in the light of its smaller CI indicative of higher specificity [21].

A comparison of circadian characteristics between drinkers and non-drinkers shows that alcohol consumption is associated with a statistically significant increase in both the MESOR and in the circadian amplitude of systolic and diastolic BP, while it does not affect pulse pressure, Fig. (9).

The BP MESOR, however, is only slightly increased by 5 to 7% (SBP: 135.5 vs. 129.1 mmHg; DBP: 81.6 vs. 76.2 mmHg). By contrast, alcohol consumption has a much larger influence on the circadian amplitude of BP, which is increased by over 20% (SBP: 27.1 vs. 22.0 mmHg; DBP: 19.1 vs. 15.4 mmHg). It is thus conceivable that in patients with a strong circadian variation at the outset, alcohol
Alcohol consumption may bring about CHAT, thereby increasing their cardiovascular disease risk. Results in Figs. (6-8) document that despite the effect of alcohol consumption on the circadian amplitude of BP, CHAT remains an independent risk factor, remaining statistically significant in the subpopulation of non-drinkers. The risk of CHAT is also independent of MH, as shown in Fig. (10). In MESOR-normotensive subjects (NT), diastolic CHAT is associated with a RR of 13.83 [0.98, 194.8] (P=0.014) among non-drinkers. Among drinkers, the only subject with diastolic CHAT suffered a morbid event, whereas none of the subjects with an acceptable circadian amplitude of diastolic BP had an event (Fisher’s exact test: P=0.026). In (treated) MESOR-hypertensive patients, diastolic CHAT is associated with a RR of 2.45 [0.89, 6.72] (P=0.108) among non-drinkers, and with a RR of 3.00 [1.47, 6.14] (P=0.007) among drinkers. The fact that numerically, cardiovascular disease risk is higher among drinkers than among non-drinkers is in keeping with the increase in the circadian amplitude of BP associated with alcohol consumption. The much larger RR observed in normotensive subjects by comparison with MESOR-hypertensive patients may be accounted for by the earlier documentation that CHAT may be a condition preceding the onset of MESOR-hypertension observed both in the laboratory [22] and in the clinic [23, 24].

**DISCUSSION AND CONCLUSION**

In another study [25], the BP of 7 clinically healthy Japanese volunteers was studied in a cross-over design. For 5 days, the subjects consumed either 40 g of alcohol by day or fruit juice, the two spans on alcohol and juice being separated by a 1-week washout. In the absence of a difference in BP MESOR, alcohol consumption was clearly associated with a large increase in overall cardiovascular disease risk among drinkers as well as non-drinkers. © Halberg (with permission).

**Fig. (10).** In both normotensive subjects and in MESOR-hypertensive patients, diastolic CHAT is associated with a large increase in overall cardiovascular disease risk among drinkers as well as non-drinkers.
when alcohol consumption was reduced from 8 to 4 and then to 2 ounces of liquor each evening, the amplitude dropping from 13.7/10.9 to 8.8/8.0 and 7.9/7.8 mmHg (systolic/diastolic BP) [27].

Our results indicate that alcohol consumption is primarily associated with an increase in the circadian amplitude of BP, with no major change in the BP MESOR. Heavy alcohol drinking is reportedly associated with an increased incidence of morbid cardiovascular events, thought to be mediated by changes in BP, yet there is no consensus on the effect of alcohol intake on BP, likely because conventional trials rely mostly on casual single measurements, ignoring the BP dynamics. An increase of the circadian amplitude of BP above an acceptable threshold, bringing about CHAT, may be a testable mechanism possibly underlying the increase in the incidence of morbid events among drinkers.

AREAS AWAITING EXPLORATION BY SCHOLARS IN NUTRITION

While the foregoing discussion revolves around evidence related to the topic of this journal, the scope of our Tokyo/Minnesota cooperation is much broader [28-32]. It touches on chronomics in two ways, also pertinent to future nutritional investigation, namely the cosmos and the glocality of its effects. In terms of time, to think globally and act locally means that time series are analyzed as a whole (globally) as well as in systematically varied consecutive shorter sections (locally) to resolve the overall time structure, keeping in mind that cycles may change their shorter sections (locally) to resolve the overall time (globally) as well as in systematically varied consecutive sections.

CONFLICT OF INTEREST

Declared none.

ACKNOWLEDGMENT

Declared none.

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Alcohol Consumption and Vascular Variability Anomalies (VVAs)


Received: January 07, 2012 Revised: January 25, 2012 Accepted: January 27, 2012

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