Circadian variability of the cardiovascular system in hypertension: Therapeutic implications
CIP-DATA KONINKLIJKE BIBLIOTHEEK DEN HAAG

Oosting, Jan
Circadian variability of the cardiovascular system in hypertension: Therapeutic implications / Jan Oosting.
Thesis Maastricht. - With ref. - With summary in Dutch.
ISBN 90-9012201-X
NUGI 743
Subject headings: chronopharmacology / autonomic nervous system regulation
Circadian variability of the cardiovascular system in hypertension: Therapeutic implications

PROEFSCHRIFT

ter verkrijging van de graad van doctor
aan de Universiteit Maastricht,
op gezag van Rector Magnificus, Prof. dr. A.C. Nieuwenhuizen Kruseman,
volgens het besluit van het College van Decanen,
in het openbaar te verdedigen,
op vrijdag 27 November 1998 op 12.00 uur

door

Jan Oosting

geboren te Zutphen in 1963
Promotor: Prof. dr. H.A.J. Struijk-Boudier
Co-promotor: Dr. B.J.A. Janssen

Beoordelingscommissie:
Prof. dr. P. de Leeuw (voorzitter)
Prof. dr. B. Lemmer (Heidelberg, Duitsland)
Prof. dr. J-L. Elghozi (Parijs, Frankrijk)
Prof. dr. D.W. Slaaf
Dr. L.L.H. Peeters

Het onderzoek beschreven in dit proefschrift is mede mogelijk gemaakt door financiële bijdragen van de Nederlands Hart Stichting (NHS 90.258) en ALZA Corporation, Palo Alto, California, USA en ALZA.

Ook wil ik graag de Dr. Saal van Zwanenberg Stichting bedanken voor de bijdrage aan het uitgeven van dit proefschrift.
Contents

Chapter 1: Introduction ................................................................. 7
Chapter 2: Hemodynamic basis of oscillations in systemic arterial pressure in conscious rats ................................................................. 29
Chapter 3: Validation of a continuous baroreceptor reflex sensitivity index calculated from spontaneous fluctuations of blood pressure and pulse interval in rats ................................................................. 47
Chapter 4: Autonomic control of ultradian and circadian rhythms of blood pressure, heart rate, and baroreflex sensitivity in SHR ............. 63
Chapter 5: Circadian and ultradian control of cardiac output in spontaneous hypertension in rats ................................................................. 81
Chapter 6: Time-Dependent Efficacy of Antihypertensive Agents in Spontaneously Hypertensive Rats ................................................................. 101
Chapter 7: Circadian-phase dependent pharmacodynamics of angiotensin converting enzyme inhibitors in spontaneously hypertensive rats ................................................................. 115
Chapter 8: General discussion .............................................................. 137
Chapter 9: Nederlandse samenvatting .................................................. 149
Dankwoord ............................................................................. 156
Curriculum Vitae ..................................................................... 157
Chapter 1
Introduction
1. Introduction

In all life forms most biological variables show different values depending on the time of day. The explanation for these circadian, or about 24 hour, differences must be sought in the adaptation to changing external stimuli throughout the day. The differences between darkness and light are possibly the strongest amongst these stimuli. Many organisms relate their level of activity to the level of light intensity. Therefore, most organisms are entrained to the 24-hour light/dark cycle.

In the course of evolution several mechanisms have developed which show endogenous circadian rhythmicity in the absence of external time clues. Molecular mechanisms have been identified that influence the circadian clock in the fungus Neurospora crassa and the fruit fly drosophila melanogaster. In Neurospora crassa the expression of the gene 'frequency' shows a circadian rhythm. When mutations in this gene appear the circadian behaviour is changed (see Figure 1.1). Both shortening and extending of the period is possible. In drosophila melanogaster a pair of proteins, timeless and per, has been shown to cause circadian behavioural rhythms which disappear when mutations in the genes of these proteins occur. These proteins are entrained to the light dark cycle because the timeless protein degrades in the presence of light.

In mammals a small part of the hypothalamus in the brain, the supra chiasmatic nucleus (SCN), has been shown to be involved in many circadian rhythms. In the

![Diagram of a cell expressing a circadian rhythm](image)

**Figure 1.1** A schematic diagram of a cell expressing a circadian rhythm. The *frq* gene encodes the FRQ protein that has multiple roles, of which the best established is to regulate the amount of *frq* mRNA as a part of a negative feedback loop that constitutes the circadian oscillator. The other is to activate clock-controlled genes (ccg's) (adapted from Aronson)
absence of external clues the SCN is able to maintain a circadian pattern. The suprachiasmatic nucleus is synchronized or entrained by the actual dark/light pattern via nervous connections from the retinal cells in the eye. Light therapy is already used to cause phase-shift in disorders like jet lag and shift work sleep disturbance. Extraocular light-sensitive sites capable of phase shifting circadian patterns have recently been shown where light pulses on the knee were able both to phase advance, and phase delay the circadian pattern of body temperature\(^6\). This method could be used to give light exposure during the night, when people are usually asleep. Other processes are able to entrain the SCN too, like feeding behaviour and physical activity. Melatonin, a hormone produced in the pineal gland, plays a key role in the genesis of several circadian rhythms. The SCN has binding sites for melatonin\(^6\), and melatonin may be used to manipulate the circadian clock, for instance to decrease to effects of jet-lag\(^9\). The presence of endogenous circadian rhythms is important for the anticipation of daily recurring activities. For instance, the organism can prepare itself for an increase in physical activity when the day starts, even when it is still dark\(^11\), in stead of only reacting to changing conditions.

The time of day has been shown to be important in several pathophysiological states. For instance, asthmatic episodes are more frequent in the night, the susceptibility of cancer cells for chemotherapy shows day-night differences. Hypertension-related events such as myocardial infarction show a peak incidence in the early morning\(^12\). In the case of asthma and cancer this has lead to changes in therapeutic regimens in order to improve pharmacological therapy\(^13\). The timing of anti-hypertensive drugs is not yet part of the World Health Organization guidelines for the management of hypertension\(^14\), although recently drug formulations have been introduced which are designed for the timed delivery of anti-hypertensive agents\(^15,16\).

2.1. Cardiovascular function

The cardiovascular system is responsible for the supply of nutrients and oxygen to different parts of the body and the discharge of waste products. The heart is the pump which provides the energy to distribute blood to all parts of the body. Blood is distributed to all organs through arteries. In capillaries nutrients and waste products are exchanged between blood and the organ, and through veins the blood is carried back to the heart.

There are a number of parameters to describe the function of the cardiovascular system. Cardiac output and total peripheral resistance determine mean arterial pressure. Cardiac output is the total blood flow from the heart and is determined by heart rate and stroke volume. The number, caliber, and geometry of small arteries and arterioles mainly determine total peripheral resistance.

In order to provide a steady flow of blood to the body blood pressure must be tightly regulated\(^17\). There are several mechanisms that control blood pressure. On the short term, in the order of seconds to minutes, the autonomic nervous system, by means of the baroreflex\(^18,20\), plays a major role in regulating arterial blood pressure. Pressure sensors in the carotid sinus and the arch of the aorta detect changes in blood pressure. Brain centres in the medulla oblongata correct these blood pressure changes
by changing heart rate and the resistance in small arteries. More recent studies suggest that nitric oxide plays a role in short-term blood pressure control as well. When blood pressure increases, shear stress in the arterial wall increases too\cite{21,22}. This in turn leads to release of nitric oxide, a potent vasodilator, thereby reducing blood pressure. On a longer time frame, of minutes to hours, humoral factors and hormones like the renin angiotensin system, vasopressin, bradykinin, and atrial natriuretic peptide play an important role\cite{24,25}. On the long term, in the order of days to weeks, blood pressure is controlled by the kidney, which keeps the amount of body fluid input and output balanced\cite{26,28}. On a timescale beyond that for fluid volume control, structural changes in the heart and in the vascular architecture play an important role in cardiovascular control\cite{29,32}.

Besides mechanisms to control blood pressure, the body also has mechanisms to control blood flow to separate organs. First there are auto-regulatory mechanisms that keep flow through organs relatively constant when pressure changes, by increasing the resistance when blood pressure rises and vice versa\cite{31}. On the other hand flow to an organ must increase when that organ has an increased metabolic demand, for instance flow to muscles may increase several times during physical exercise\cite{32}. The exact mechanisms that control local organ flow are a topic of on-going research, but the myogenic response and endothelium derived vaso-active substances like nitric oxide, endothelin and prostacyclin seem to play an important role\cite{33}.

The mechanisms described here interact on several levels\cite{34,35,36,37}. Therefore it can be difficult to differentiate between the effects of different mechanisms, especially in whole animal preparations or human subjects. Blockade, or stimulation, of one mechanism is bound to be followed by reactions of another regulatory mechanism. The redundancy of control mechanisms for blood pressure is a sign for the importance of blood pressure control for the organism.

2.2. Variability in cardiovascular parameters

A considerable variability exists in the core parameters of cardiovascular function\cite{39,40}. Variability can be caused by internal sources, like the control mechanisms regulating cardiovascular function, or circadian rhythms, or by external sources, like positional changes or cardiovascular active drugs. A part of the variability can be attributed to regular oscillations. These oscillations are divided into categories, dependent upon their period. Ultradian rhythms have a period shorter than a day. Circadian rhythms have a period of about one full day, and infradian rhythms have periods of more than one day.

2.2.1. Ultradian rhythmicity in cardiovascular function

When the frequency content of cardiovascular parameters is examined by spectral analysis, a few frequencies are predominantly present\cite{35,41}. If the spectral power in a frequency band is caused exclusively by a certain regulatory mechanism, the technique of spectral decomposition of time signals can be used to study that mechanism without having to isolate a part of an organism. Therefore a lot of studies have been devoted to
assign physiological mechanisms to certain frequencies. The peaks most commonly studied with spectral analysis are those that are related to actions of the autonomic nervous system. These are normally divided in a high frequency range and a low frequency range. The parasympathetic nervous system can react fast enough on pressure fluctuations caused by respiration and is usually responsible for the fastest or high frequency oscillations in heart rate (around 0.25-0.3 Hz in humans, and about 1.2 Hz in rats). There is still debate on the origin of low frequency oscillations in blood pressure (< 0.1 Hz in humans, about 0.4 Hz in rats). These could be caused by the sympathetic nervous system, or by a combination of parasympathetic and sympathetic influences. Local regulatory systems, and mechanical factors like breathing, may interact with arterial pressure fluctuations in this frequency range.

Sometimes also even slower oscillations of blood pressure are identified of 26 minutes, or with periods between 1 and 2 hours, but in general, an inverse linear relationship can be found between the logarithm of the frequency and logarithm of spectral power of blood pressure and heart rate in this lower frequency range. This so called 1/f characteristic indicates the absence of a single dominant regulatory mechanism and is suggestive of non-linear system characteristics with multiple regulatory processes acting on overlapping time scales.

2.2.2. Circadian rhythmicity in cardiovascular function

All hemodynamic variables show differences between day and night. For heart rate and blood pressure, which are relatively easily measured, it has long been known that they are higher during the light period in diurnal animals (and humans) and that they are higher during the dark period in nocturnal animals. The circadian rhythm of blood pressure or heart rate has not yet been shown to persist under constant light conditions in humans, but rats show a circadian period of blood pressure and heart rate of longer than 24 hours in constant darkness. The daily variations in heart rate and blood pressure are determined by physical activity mostly. In shift-workers the circadian pattern of blood pressure is adapted to the working pattern. However, subjects that are kept supine continuously, still exhibit circadian variations of these parameters, showing the prevalence of an endogenous rhythm. During 6 weeks of head-down tilt the circadian variation of diastolic blood pressure disappears almost completely within a few days, the circadian variation of systolic blood pressure is halved, while the circadian variation of heart rate is not affected. Cardiac output and stroke volume also exhibit a circadian rhythm with higher values during the active period. Because cardiac output decreases relatively more at night than blood pressure, total peripheral resistance is higher during the resting period. The circadian variation in vascular resistance of individual organs may differ between organs because the relative flow distribution varies throughout the day. For instance in normotensive humans, forearm peripheral resistance is higher during the active phase of the day, whereas total peripheral resistance is estimated to be lower at that time. These differential distributions can be explained by the sympathetic control of muscle vascular beds.

Regulatory mechanisms important for cardiovascular function show oscillatory behaviour too. In humans renin is rhythmically released during sleep. Circadian vari-
ations are also seen. The balance between the sympathetic and the parasympathetic part of the autonomic nervous system is shifted to the sympathetic part during the active period of the day when blood pressure and heart rate are high, whereas the parasympathetic part is dominant during the resting period of the day. Circadian patterns of plasma noradrenaline are synchronous to the blood pressure pattern in humans and rats. The gain of the baroreflex is higher during the resting period. The hormones with cardiovascular pressor effects show a circadian rhythm that is opposite to the blood pressure profile. In humans the renin angiotensin system, vasopressin, and cortisol all have their highest activity during the night and early morning. In renin transgenic rats (TGR) even an inverted circadian blood pressure pattern has been observed. The high levels of these hormones during inactivity could be especially important in order to retain fluid during the resting period, and keep diuresis at a low level. Endocrine mechanisms that lower blood pressure are generally more active during the resting phase, and these mechanisms are therefore synchronized to the circadian rhythm of blood pressure. Atrial natriuretic peptide shows circadian rhythms with higher values during the resting phase. Nitric oxide guanylyl cyclase expresses higher activity in the aorta, and administration of L-NAME produces larger increases in blood pressure during the resting period in rats, showing that the blood pressure lowering effect of nitric oxide is synchronized to the blood pressure pattern.

The excretion of fluid and minerals by the kidney shows a strong circadian rhythm. The excretion of water and sodium, and the ingestion of water is highest in the active period, and lower during sleep. In addition to blood pressure, the excretion of fluid by the kidney is influenced by a large number of factors, like the sympathetic nervous system, the renin-angiotensin system, aldosterone, and vasopressin. These systems all have their own circadian rhythm as well. It is therefore difficult to obtain information on the circadian variability of the direct influence of pressure on renal fluid excretion. In rats it has been shown that urinary excretion of water, sodium and potassium show a circadian rhythm both in blind rats and in rats under constant illumination, furthermore it lasts about a week before the circadian pattern of excretion is synchronized to a phase-shift of 12 hours of the light-dark pattern.

2.2.3. Infrahadian rhythmicity

A number of rhythms have periods longer than a day. The most important of these are weekly, about monthly, and yearly rhythms. Weekly or circaseptan rhythms can be attributed to the human inclination to have one sleep out, a tendency to delay bedtime in the weekend, and a weekly working schedule. This causes a phase delay of the circadian system of about half an hour. On the other hand, circaseptan rhythms in sodium excretion have been found in laboratory animals that were on a daily pattern without changes in the weekend. The risk of myocardial infarction is also unevenly distributed throughout the week. The incidence is highest on Monday morning, especially in the working population. Monthly rhythms in cardiovascular function can be observed predominantly in women. Within the menstrual cycle large variations of oestrogen occur, a hormone with important cardiovascular properties. A yearly cycle has been found in the occurrence of angina pectoris. This effect is stronger in regions with clear climatic differences between the seasons, since this phenomenon seems to be
mainly caused by differences in temperature between the seasons.92-94

3.1. Hypertension

Among the human population a considerable variation exists in the level of arterial blood pressure. Only at relatively high levels blood pressure in itself causes symptoms like head ache and retinal bleeding. Prospective epidemiological research has shown that blood pressure is positively related to the occurrence of cardiovascular disease,95,96 and it is therefore that high blood pressure is treated, even at levels at which no acute symptoms occur.

Hypertension, defined as a resting diastolic blood pressure of more than 95 mmHg and/or a systolic blood pressure above 160 mmHg,97 is very common in westernised countries, and cardiovascular disease is the most common cause of death. Through an increased mechanical load on the heart and the arterial side of the vascular system hypertension causes left ventricular and arterial hypertrophy, and is a main factor in the pathogenesis of atherosclerosis. Coronary atherosclerosis again is the main cause of myocardial infarction. High blood pressure itself is the main risk factor for stroke and renal failure.97

A cause for hypertension can be found in about 10% of cases. Common causes for these secondary forms of hypertension are adrenal, renal, or renovascular disease, or drug related causes. This means that in about 90% of cases the aetiology of hypertension is unknown. This most common form of hypertension is called primary or essential hypertension. Although the actual cause of essential hypertension is unknown, a number of risk factors have been identified. Genetic predisposition, environmental factors (i.e. high salt intake and high alcohol consumption), and psychological stress have all been shown to increase the risk for hypertension.97

From a hemodynamic point of view hypertension is characterized by high blood pressure, high total peripheral resistance but normal cardiac output, heart rate and stroke volume.98 There are several theories on the primary defect to explain the increase of total peripheral resistance in hypertension.

The first theory was formulated by Borst & Borst-de Geus and Guytor and puts emphasis on the kidney.99,100 Improper handling of volume and/or sodium leads to overfilling of the vascular system. This high intravascular volume leads to an increase in cardiac output. Autoregulatory mechanisms in the organs prevent over perfusion by increasing their resistance resulting in increased blood pressure. Because the kidney excretes more water and sodium when blood pressure is high, the balance between fluid input and output is restored at the expense of a higher blood pressure.

The second theory proposes increased sensitivity of the vascular system to vasoconstricting agents with associated structural changes of the resistance arteries as the primary defect.101-103 Hypertrophied vessels are hypersensitive to vasoconstricting agents and this leads to a form of positive feedback whenever less pressure increasing influence is capable of increasing blood pressure more.

More recently a third theory has been postulated that attributes the increase in total peripheral resistance to a decreased number of vessels in the microcirculation. Both regression of existing vessels, and insufficient growth of vessels during develop-
3.2. Circadian aspects of hypertension and cardiovascular risk

The development of devices to automate blood pressure measurements has made it possible to study circadian aspects of hypertension. The use of these devices has identified a form of hypertension that is characterized by an increase of blood pressure by psychologically stressful events. The blood pressure pattern during normal daily activities is comparable to normotensive subjects, but for instance during the interview, and examination by a doctor blood pressure rises. This so-called 'white coat hypertension' turns out to be rather benign in nature.

In essential hypertension the circadian variability of hemodynamic parameters is largely preserved. However there is a subset of the hypertensive population in which blood pressure does not fall during sleep, the so-called non-dippers. Also in secondary forms of hypertension the difference between day and night blood pressure is usually small. Non-dippers have an increased morbidity of cardiovascular disease and end organ damage, like left ventricular hypertrophy and retinopathy, compared to hypertensive subjects that do show a fall in blood pressure during sleep. It is not yet clear what the causal relation is in this observation. It can be either that the non-dipping phenomenon is harmful, or that end-organ damage leads to non-dipping. Extreme dipping may predispose to brain damage in elderly hypertensive patients, and thus non-dipping may be a protective reaction of the body to preserve perfusion pressure to the brain. Alternatively non-dipping may be caused by the increased importance of the renin-angiotensin system in some forms of hypertension with its higher pressor activity during sleep. The circadian variability of heart rate and cardiac output is hardly affected by hypertension.

The incidence of cardiovascular events like myocardial infarction, stroke, and sudden death shows a circadian variation too. These events occur most often in the early morning. A number of physiological processes have a circadian rhythm that can contribute to the increased incidence of cardiovascular events in this period. Blood pressure and heart rate rise in this period of the day. Together with an increase in blood pressure variability through a decrease in baroreflex function this imposes an increase in mechanical stress to the vascular wall, and makes atherosclerotic plaques more susceptible to rupture. Platelet aggregability peaks in this period too. A concurrent decrease in the counteracting activity of plasminogen activator makes the formation of a blood clot more easy. Cortisol shows peak levels during the early morning, increasing the sensitivity of the coronary arteries for the vasoconstrictive properties of the catecholamines, which show an increase in plasma levels at this period too.

Although most evidence supports the fact that the patterns of these physiological processes are caused by the increase in physical activity during the early morning, it cannot be ruled out that these rhythms are partly endogenous. The uncertainty
about the origin of circadian patterns is one of the fundamental topics in chronobiologic research. Most circadian patterns are composed of an endogenous component, driven by an internal oscillator, and so-called masking effects, direct reactions to changes in the environment. The combination of these effects is probably beneficial to the individual. The endogenous rhythm can adapt the organism to predictable changes in the environment, while the masking effects react to unpredictable changes.

4.1. Treatment of hypertension: risk reduction

Meta-analysis of several interventional trials has shown that treatment of moderate hypertension decreases the incidence of both stroke and coronary heart disease. The reduction in stroke is much more pronounced than the reduction in coronary heart disease, and the morbidity of cardiovascular disease is still higher in treated hypertensive subjects, than it is in normotensive subjects. Explanations for the partial failure of anti-hypertensive treatment to reduce cardiovascular morbidity are 1) Blood pressure is still higher in treated hypertensive subjects, than in the normotensive population. This can be caused by non-compliance to therapy. 2) The anti-hypertensive agents used in these studies were mainly diuretics and beta blocking drugs which can have adverse effects on lipid and glucose metabolism. More modern drugs like calcium antagonists or ACE-inhibitors do not have this adverse effect, or even improve glucose and lipid metabolism. Data of prospective studies of these drugs on mortality is not yet available. 3) Structural damage or changes like left ventricular hypertrophy and arterial stiffening by hypertrophy or atherosclerosis in the period before treatment may be irreversible. 4) Other risk factors contributing to (the development of) hypertension or the risk for cardiovascular morbidity, like smoking, high salt intake, and high cholesterol level, may still be present during pharmacological treatment.

It appears important that hypertensive subjects should be treated so that diastolic blood pressure is between 85 and 90 mmHg. It is reported that reduction of blood pressure below this value increases cardiovascular morbidity again, the so-called J-shaped phenomenon. It seems that in the untreated population this J-shaped phenomenon does not occur and the morbidity from cardiovascular disease levels out below 90 mmHg, while in treated hypertension morbidity increases when blood pressure is lowered below 85 mmHg. The explanation for the J-shape could be that perfusion of vital organs is compromised by low pressure, because of atherosclerosis or hypertrophy of the arterial system. The term essential hypertension is appropriate in these cases, because high blood pressure is a life saving adaptation of the body.

In recent years some hypertension management strategies have been developed which are focussed more toward the individual patient. Therapy is initiated and evaluated dependent on absolute risk for cardiovascular morbidity, by the presence of risk factors and established end-organ damage.

4.2. Treatment of hypertension in relation to time of day

The realization that cardiovascular incidents are most prominent in the early
morning has generated interest in the amount of protection that anti-hypertensive drugs give at this period of the day\textsuperscript{147, 148}.

Using ambulatory blood pressure devices many studies have been carried out with various classes of anti-hypertensive drugs to assess the effects on the 24-hour blood pressure pattern. The general conclusion of these studies is that anti-hypertensive drugs with a sufficient long half-life all lower blood pressure throughout the day and night. The circadian pattern of blood pressure during treatment resembles the pattern during placebo, although at a lower level. A meta-analysis to compare reduction of blood pressure during the day and night-time showed no difference for beta blockers and ACE-inhibitors, and with calcium antagonists blood pressure was 2 3/4 more decreased during the day-time\textsuperscript{149}(Table 1.1).

<table>
<thead>
<tr>
<th>Class of drug</th>
<th>Studies</th>
<th>N</th>
<th>SBP\textsubscript{Day} (mmHg)</th>
<th>DBP\textsubscript{Day} (mmHg)</th>
<th>SBP\textsubscript{Night} (mmHg)</th>
<th>DBP\textsubscript{Night} (mmHg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beta-blockers (Once)</td>
<td>9</td>
<td>153</td>
<td>9.8</td>
<td>8.6</td>
<td>13.0</td>
<td>10.7</td>
</tr>
<tr>
<td>Beta-blockers (2-3x)</td>
<td>5</td>
<td>67</td>
<td>13.1</td>
<td>10.6</td>
<td>15.8</td>
<td>13.8</td>
</tr>
<tr>
<td>Calcium antagonists (once)</td>
<td>14</td>
<td>213</td>
<td>10.6</td>
<td>6.2</td>
<td>10.8</td>
<td>9.3</td>
</tr>
<tr>
<td>Calcium antagonists (2-3x)</td>
<td>11</td>
<td>164</td>
<td>11.5</td>
<td>11.1</td>
<td>12.7</td>
<td>10.1</td>
</tr>
<tr>
<td>ACE inhibitors (once)</td>
<td>8</td>
<td>104</td>
<td>8.9</td>
<td>7.8</td>
<td>11.1</td>
<td>8.7</td>
</tr>
<tr>
<td>ACE inhibitors (2-3x)</td>
<td>4</td>
<td>54</td>
<td>11.1</td>
<td>10.1</td>
<td>12.0</td>
<td>11.7</td>
</tr>
</tbody>
</table>

*Once: Once daily dosing; 2-3x: 2-3 times daily dosing; Studies: number of studies in this class; N: total number of subjects. Values are weighted averages.*

The United States Food and Drug Administration has introduced guidelines for the registration of new anti-hypertensive drugs which require a minimum trough-to-peak ratio\textsuperscript{150-153}. The trough-to-peak ratio is calculated by dividing the remainder of the blood pressure lowering effect just before a new dose by the maximum blood pressure lowering effect of a drug (Figure 1.2). The trough-to-peak ratio was designed to ensure efficacy at the end of the dosing interval, without having to resort to inappropriately high doses. Trough-to-peak ratio is dependent on factors like elimination half-time, type of elimination, pharmacokinetic-dynamic properties, dose, and dosing interval\textsuperscript{153}. There has been considerable discussion on the implementation and value of the trough-to-peak ratio\textsuperscript{154, 155}, also because it proves difficult to determine the peak effect of drugs in relation to the dosing time and plasma concentrations of drugs\textsuperscript{156}. The frequency of blood pressure measurements must be high enough to detect the peak effect reliably\textsuperscript{153}.

The fact that blood pressure and heart rate pattern are comparable in treated and non-treated subjects is rather surprising since there are numerous factors which can produce differences in the efficacy of these drugs throughout the day. Pharmacokinetics may vary throughout the day by variations in absorption, distribution, or clear-
Figure 1.2 Trough peak ratio For the calculation of trough-to-peak ratio treatment effect is important. The trough value is the lowest difference between control blood pressure and blood pressure during treatment, which is usually in the period just before the new dose is taken. The peak value is taken at the time of the maximum blood pressure lowering effect. This time point is dependent on the drug and formulation used. The measurement of blood pressure must be sufficiently often in order to ensure that the maximum blood pressure lowering effect is really found.

Furthermore, the responsiveness of effector organs or receptor sensitivity to drugs may differ during the day. Lastly, the physiological response to the changes induced by the drugs may show day-night differences.

The development of the trough-to-peak ratio has focussed attention on a flat anti-hypertensive profile of drugs, but questions remain about the desirability of such a profile. It could be advantageous to specifically target treatment to the high-risk period of the early morning besides being effective for 24 hours.

Current objectives for pharmacological anti-hypertensive therapy include 1) 24-hour control of blood pressure, also during the early morning rise in blood pressure. 2) A once daily dosing regimen to improve compliance to therapy. 3) Absence of drug related side effects; since hypertension is not a disease itself, it merely increases the risk for cardiovascular morbidity in the future. There is still room for improvement of anti-hypertensive therapy. The uneven daily distribution of cardiovascular events related to hypertension makes this a target in therapy that has not been explored fully.

5. Use of animal models to study circadian rhythms in essential hypertension

Animal research has a number of advantages over research in human subjects in the study of hypertension. Small size and a short life span make it easier to study the
natural history and pathophysiological alterations of disease. Furthermore it is possible to use interventions and experimental determinations that are not possible in humans. The genetic homogeneity and experimental control over the environment make it possible to obtain high-grade results.

In order to translate findings from animal research to the human situation, care must be taken that the studied conditions or disease have comparable effects in the animal and the human population.

A number of animal models have been designed to study hypertension. These models involve a range of experimental procedures, such as (in)breeding, surgical interventions, genetic alterations, drug or hormone administration or exposure to noxious stimuli. The spontaneously hypertensive rat (SHR) has been extensively used as a model for essential hypertension. The SHR was originally developed by inbreeding in 1963. Hemodynamics, vascular reactivity, sympathetic nerve activity, and levels of cardiovascular active hormones have all been shown to show a comparable effect in human essential hypertension and SHR. Goldblatt first introduced the 2-kidney 1-clip model. During surgery the lumen of one renal artery is narrowed. This causes a lower pressure in the affected kidney, which secretes more renin in order to normalize pressure in that kidney. The model reflects hypertension as seen during renal artery stenosis in humans. In 1990 a new model of secondary hypertension was developed by the introduction of the mouse Ren-2 salivary gland renin gene into the genome of the rat. In these rats very high levels of renin are secreted, resulting in markedly elevated blood pressure levels.

Recent advances in genetic engineering in mice have also produced models of hypertension, especially with transgenic animals with human renin or angiotensin genes. However because of the size of these animals long-term blood pressure measurements are not feasible yet.

In SHR and 2K-1C circadian variability of blood pressure been shown to exhibit blood pressure patterns which are comparable with those in human hypertension, although with a phase difference of about 12 hours because of the nocturnal nature of these animals, while Ren-2 TGR show an inverted circadian blood pressure pattern.

In the present thesis SHR are used because it serves as a model of human hypertension in aspects under investigation in the studies. The neurohumoral status, the circadian variability of hemodynamic variables, and the response to anti-hypertensive drugs is comparable to the effects seen in essential hypertension in humans.

6. Aim of this thesis

From the introduction it can be concluded that anti-hypertensive therapy can be improved. The uneven distribution of incidence of cardiovascular disease over the day makes this a target in therapy which has not been explored fully. In order to make fundamental changes in therapeutic strategies it is important to acquire knowledge of the basic mechanisms that are responsible for blood pressure control throughout the day. The autonomic nervous system seems to play a key part in the circadian variability of hemodynamic parameters. Levels of sympathetic activity and plasma levels of norepinephrine are synchronous to the blood pressure, and heart rate pattern. The chapters
2-5 of the thesis are aimed at understanding the role of the autonomic nervous system in the regulation of short-, and long-term rhythms of blood pressure, heart rate, and cardiac output in hypertension. The second part of the thesis is devoted to time-dependent treatment of hypertension (chapters 6 and 7).

The short-term variability of blood pressure has already been well established. In chapter 2 we have studied the short-term rhythms of regional and systemic blood flow, and the interaction between regional and systemic flow and blood pressure on this time scale.

Baroreflex function is one of the regulatory mechanisms of the autonomic nervous system. In humans and cats methods have been developed to measure baroreflex gain continuously from spontaneously occurring fluctuations in blood pressure and heart rate 132-134. In chapter 3 we have developed a method in rats to determine baroreflex sensitivity from spontaneous changes in blood pressure and heart rate, in order to obtain this measure continuously over 24 hours.

In hypertension the gain of the baroreflex is generally decreased 135. In order to evaluate the effects of hypertension on 24-hour baroreflex control, baroreflex function was measured for 24 hours in normotensive and spontaneously hypertensive rats in chapter 4.

In chapter 5, the relationship between cardiac output, blood pressure, heart rate, and total peripheral resistance was investigated over ultradian and 24-hour periods. Normotensive and hypertensive animals were compared under control conditions as well as during autonomic blockade.

A number of physiological processes have a circadian rhythm that increases the risk for cardiovascular events like myocardial infarction or stroke in the early morning. The accumulation of these risks at this period of the day may be synergistic. We hypothesize that it could be beneficial to desynchronize the circadian patterns of these processes. The study in chapter 6 was aimed at providing an insight into the time-dependent effects of different classes of anti-hypertensive drugs. Chapter 7 presents the effects on circadian patterns of blood pressure in hypertensive rats by repeated, timed infusion of anti-hypertensive drugs affecting the renin angiotensin system during the transition from the sleeping to the active period.

Finally, chapter 8 is aimed at an integrative discussion of all studies presented in this thesis, with the implications towards anti-hypertensive treatment.

References

Introduccion


Chapter 1


Introduction


Chapter 2

Hemodynamic basis of oscillations in systemic arterial pressure in conscious rats

Ben J.A. Janssen
Jan Oosting
Dick W. Slaaf, Department of Biophysics, University of Maastricht
Pontus B. Persson, Department of Physiology, Humboldt University (Charité), Berlin, Germany.
Harry A.J. Struijker-Boudier

Published in American Journal of Physiology (Heart Circulation Physiology 38) 1995; 269, H62-H71
Chapter 2

Abstract

In conscious resting rats, beat-to-beat fluctuations in systemic mean arterial pressure (MAP) were compared with those in cardiac output and those in blood flow in the renal, mesenteric, and hindquarter vascular beds. Spontaneous oscillations (lability) in MAP were observed in frequency bands centered about 1.6 Hz (high: HF), 0.4 Hz (mid: MF), and 0.13 Hz (low: LF). Lability of MAP was confined within the LF (~8 s) band. Lability of cardiac output, on the other hand, showed primary HF oscillations. LF oscillations in regional blood flow were most prominent in the mesenteric and renal vascular beds. In these beds, LF oscillations in blood flow showed negative phase angles with MAP, whereas those between MAP and hindquarter blood flow were positive. Cross correlation analysis indicated that ~2 s following a LF change in MAP, LF changes in mesenteric and renal blood flow occurred opposite to those of MAP. Changes in hindquarter flow were negatively correlated with those in MAP about zero time delay. Admittance gains were 1 across all frequencies for all vascular beds, indicating the absence of autoregulation. This hemodynamic pattern suggests that myogenic mechanisms predominantly control mesenteric and renal blood flow in a nonautoregulatory but rather superregulatory manner, while autonomic mechanisms regulate hindquarter blood flow. Thus, in conscious resting rats, spontaneous fluctuations in systemic arterial pressure predominantly exhibit slow (~8 s) oscillations, which do not arise from fluctuations in cardiac output, but originate from regionally specific myogenic oscillatory mechanisms contributing to resistance to flow.
Introduction

Indexes of blood pressure and heart rate variability are increasingly explored as prognostic tools in various circulatory pathophysiological states \(^{1-4}\). However, the hemodynamic mechanisms underlying fluctuations in blood pressure and heart rate, especially those that are slower than the cardiac and respiratory cycle (Traube-Hering waves), have not been fully characterized. These so-called Mayer waves or third-order waves were first described more than a century ago \(^1\). They are centered about 0.1 Hz in humans \(^{2-4,6}\), dogs \(^1\), and cats \(^4\), and about 0.4 Hz in rats \(^6,14\). After comparison between patient groups \(^{12}\) and after using various experimental interventions \(^{2,11,14}\), oscillations in these frequency bands were shown to depend largely on the integrity of the autonomic nervous system. They are possibly caused by modulation of vascular tone or heart rate either directly by a central oscillator \(^{16}\), or by resonance of the baroreflex or respiratory mediated changes in autonomic nervous activity \(^4,6,14\).

The relevance of the Mayer waves is not clear. Changes in their amplitude have been used to estimate changes in autonomic circulatory control in various diseases \(^1,3\). In addition, open-heart patients who showed vasomotor waves during cardiopulmonary bypass, needed less perioperative and postoperative inotropic support \(^17\). A considerable amount of variability in spontaneous blood pressure fluctuations may be found in frequencies below those of Mayer waves \(^1,19\). The origin of such blood pressure fluctuations, with periods between 10 s and 24 h, has not been studied in detail for several reasons. First, most studies restricted their interest to the respiratory and Mayer waves, since the resolution of data sampling or processing was too low to clearly identify oscillations at lower frequencies. Second, in conscious animal experiments, the stationarity of the blood pressure signal may be disturbed by large fluctuations due to movement \(^19\). Third, because linear logistic relations were found between spectral power and frequency \(^20,21\), deterministic nonlinear processes may underlie these blood pressure fluctuations. Consequently, independent low-frequency oscillations are not required. Nevertheless slow nonbehavioral endogenous rhythmic processes influencing arterial pressure cannot be excluded because low-frequency spectral power of blood pressure often changes after pharmacological or surgical manipulation of circulatory-controlling mechanisms \(^8,12,14,18\).

In pilot studies we tested for the presence of oscillations in arterial blood pressure in conscious rats with frequencies lower than those of Mayer waves (0.4 Hz). We were able to identify major oscillations in arterial pressure with a period of roughly 8 s, especially when rats were asleep or resting. The present study describes the hemodynamic characterization of these blood pressure rhythms. In two groups of rats we compared beat-to-beat variations in cardiac output with those occurring in blood flow in three major peripheral vascular beds. The results indicate that in resting rats the 8 s rhythm in blood pressure is not dependent of fluctuations in cardiac output but primarily caused by myogenic oscillations in renal and especially in mesenteric blood flow.
Chapter 2

Methods

Animal Preparation

Experiments were performed in male adult Wistar rats purchased from Winkelmann (Borchen, Germany). Rats were kept on a 12:12 h light-dark cycle and had free access to water and standard food pellets (type RMM-TM, Hope Farms, Woerden, The Netherlands). The experimental protocol was approved by the Institutional Animal Care and Use Committee of the University of Limburg, Maastricht.

Instrumentation of flow probes was performed under pentobarbital sodium anesthesia (60 mg/kg). For recording of central hemodynamics, electromagnetic flow probes (2.7 mm diam; Skalar, Delft, The Netherlands) were implanted on the ascending aorta. For recording of regional hemodynamics, miniaturized 20-MHz pulsed Doppler flow probes (Crystal Biotech, Holliston, MA) were implanted on the left renal artery and superior mesenteric artery and the abdominal aorta to measure renal (RBF), mesenteric (MBF), and hindquarter blood flow (HBF), respectively. Both surgical procedures have previously been described by us in detail.22 Rats were allowed to recover for 4-5 days to regain their preoperative body weight. Two days before the actual experiments, under ether anesthesia, a catheter (PE-10 heat sealed to PE-50) was inserted into the abdominal aorta via the left femoral artery for blood pressure recordings. The catheter was tunneled subcutaneously to the neck, where it was exteriorized, filled with saline, and closed with a metal plug. Experiments were completed in nine rats instrumented for recording of central hemodynamics [group I, avg body wt 290 ± 13 (SD) g] and in eight rats prepared for the regional blood flow measurements [group II, avg body wt 288 ± 11 g].

Protocol

On the day of the experiment rats were placed in experimental cages (22 x 13 x 13 cm) in a quiet illuminated room. The rats were connected to the equipment and allowed to become accustomed to the environment for 1-1.5 h. In group I, arterial pressure and ascending aortic flow was measured. In group II, arterial pressure and RBF, MBF, and HBF were measured. Recordings were made for 30 min during the resting or sleeping state to minimize locomotor-induced changes in hemodynamics. During the recording period behavioral activity of the rats was scored every 15 s using a scale of 1-5 corresponding to the following states: 1) sleeping; 2) awake but not moving; 3) small movements such as making a turn before lying down again; 4) vigorous movements associated with grooming or exploring; and 5) standing against the wall of the cage. The average behavioral activity in groups I and II was 1.2 ± 0.2 (SD) and 1.3 ± 0.2, respectively. In both groups, the time span in which the average index was > 2 was < 5% of the whole recording period. For this reason we did not exclude periods with higher behavioral activity from the analyses. Breathing rhythm was assessed by watching the abdominal movements every 10 min. Respiratory frequencies were not different between groups [group I: 1.72 ± 0.33 (SD), group II: 1.58 ± 0.19].

Data acquisition

The arterial catheter was connected to a low-volume pressure transducer (Micro Switch 156PC15GWL, Honeywell, Amsterdam, The Netherlands) for the measurement
of arterial blood pressure. Cardiac output was measured as ascending aortic flow by using a sine-wave electromagnetic flowmeter (model 1401, Skalar). Late diastolic flow was taken to be zero. Regional blood flow velocities were measured as Doppler shift (in kHz) with a three-channel 20-MHz pulsed ultrasound Doppler flowmeter (Bioengineering Department, University of Iowa, Iowa City, IA). Zero flows were obtained electronically. Arterial pressure and flow signals were recorded on a beat-to-beat basis with the aid of a data-acquisition program developed by the Engineering Department of the University of Limburg, Maastricht, The Netherlands. All signals were sampled continuously with a minimum frequency of 500 Hz using a DT 2814 A/D converter board (Data Translation, CN Rood, Rijswijk, The Netherlands) housed in a 486 IBM-compatible computer. The time lag between the beginning of the ascending aortic flow and the abdominal aortic pressure wave was ~20 ms. The time lag between the beginning of the regional flow waves and the abdominal pressure wave was ~6 ms. Beat-to-beat values of mean arterial pressure (MAP), pulse interval (PI), stroke volume (SV), and cardiac output (CO) were calculated on-line using an algorithm that used the end-diastolic value of the aortic (group I) and HBF signal (group II) to determine the interbeat interval. Total peripheral resistance (TPR) was estimated by dividing MAP by CO. All beat-to-beat variables were stored on disk for off-line analysis. Because of technical problems, the MBF of one rat could not be recorded.

**Data processing**

The beat-to-beat signals of arterial pressure and central or peripheral hemodynamics were replayed on screen for visual inspection. Data segments with artifacts due to obstruction of the arterial catheter or electrical noise were eliminated (<0.2% of all beats). For all variables means ± SD and coefficient of variation were determined. Furthermore, as an index for short-term lability, the average relative (as percentage of mean) beat-to-beat differences of all signals were calculated.

The dynamic interactions of arterial pressure and blood flow fluctuations were investigated in the frequency domain using a fast Fourier transform computer program developed at the Department of Physiology, University of Heidelberg, Germany. For this analysis technique equidistantly sampled data are required. Therefore the beat-to-beat data points were converted into equidistant time series using an algorithm that extracted from sequential 230-ms time windows the maximum value of each parameter. Because the interbeat interval was never lower than 226 ms during the recordings, the width of the window was set at 230 ms to avoid aliasing. The resulting (4.35 Hz) time series were divided into half-overlapping sequential sets of 512 data points (each ~100 s). Before calculation of spectral density power, each segment was subjected to linear trend removal and cosine tapering of the first and last 60 data points. Spectral density power was expressed relative to total power obtained by integrating all values from 0 to 2.175 Hz (Nyquist frequency).

In addition to the spectral density power, the transfer function was calculated, assuming MAP as input and blood flow as output, as previously described in detail. For each segment of 512 data points, the magnitude and phase of the transfer function was calculated as the quotient of the input and cross spectrum. Variance was reduced by averaging the resulting sequential data sets. The admittance gain was calculated by normalizing the magnitude of the transfer function by the average con-
Figure 2.1. A: tracings from rat 1 showing example of 512 beats recordings of pulse interval (PI), mean arterial pressure (MAP), cardiac output (CO), stroke volume (SV), and total peripheral resistance (TPR), respectively, in a conscious resting rat (Group I). B: Spectral power plots of corresponding tracings. Low (LF: 0.09-0.19 Hz)-, mid (MF: 0.30-0.50 Hz)-, and high (HF: 1.20-2.00 Hz)-frequency bands are indicated by black bars.

Figure 2.2. A: tracings from rat 5 showing example of 512 beats recordings of PI, MAP, and renal (RBF), mesenteric (MBF), and hindquarter (HBF) blood flow in a conscious resting rat (Group II). B: Spectral power plots of corresponding tracings. The LF, MF, and HF bands are indicated by black bars.

Admittance calculated as the ratio of the average blood flow and pressure in the whole 30-min time series. The admittance gain is the ratio of the fractional variation in flow and pressure. A value of 1 for the gain indicates that when the input (MAP) fluctuates by some fraction of the mean value, the output (flow) undergoes the same fractional variation, as would be the case in a rigid vascular bed. A value of < 1 signifies that the fractional variation in blood flow was smaller than in blood pressure. This would indicate autoregulatory behavior. When the admittance gain is > 1, the pressure oscillations are then amplified in the heart or peripheral organs. Alternatively, these admittance values may indicate that the elevated fractional variations in blood flow are caused by autonomous fluctuations originating in the heart or vascular bed and are reflected in the blood pressure signal. The phase indicates the temporal relationship between the blood pressure and blood flow signals in the frequency domain, whereas the cross-correlation coefficient indicates their correlation in the time domain. In addi-
Hemodynamic basis of blood pressure oscillations

Figure 2.3. Comparison of average spectral power density (solid line) ± SE (dotted line) in groups I (PI, MAP, CO, SV, and TPR, A, n=9) and II (PI, MAP, RBF, MBF, HBF, B, n=7-8). For each rat data were averaged over ~32 overlapping sequential 100-s periods. LF, MF, and HF frequency bands are indicated by black bars.

Figure 2.4. Average admittance gain, phase delay, and coherence (significant when >0.4, p < 0.05) of MAP vs. CO. Data are means ± SE of 9 rats. For each rat data were averaged over ~32 overlapping sequential 100-s periods. LF, MF, and HF bands are indicated by black bars.

In addition to these calculations, the coherence function was also computed. Coherence can have a value between 0 and 1 and is a frequency domain estimate of the degree that the flow variance can be explained by a linear operation of the blood pressure variance. The coherence is 1 if no other variables than blood pressure affect the flow oscillations. Finally, for each rat, the results obtained from these analysis techniques were averaged over the sequential data segments to reduce the variance.

The spectra were divided into four different frequency domains: 1) a high-frequency range (HF: 1.2–1.8 Hz) containing oscillations that are linked to respiration; 2) a midfrequency range (MF: 0.3–0.6 Hz) containing oscillations that may stem from sympathetic and parasympathetic influences\(^{26,27}\); 3) a low-frequency range (LF: 0.08–0.18 Hz) representing oscillations that are thought to result from vascular myogenic responses\(^{26,27}\); and 4) a very low frequency range (VLF: 0.03–0.08 Hz) that comprises all undefined lower frequency oscillations but may include a tubuloglomerular feedback induced oscillation as has been described for RBF in anesthetized rats\(^{26,27}\). Note that the
Table 2.1. Average values of hemodynamic parameters in groups I (central hemodynamics) and II (regional hemodynamics)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Group I</th>
<th>Group II</th>
</tr>
</thead>
<tbody>
<tr>
<td>MAP, mmHg</td>
<td>9</td>
<td>8</td>
</tr>
<tr>
<td>PI, ms</td>
<td>9</td>
<td>8</td>
</tr>
<tr>
<td>CO, ml/min</td>
<td>9</td>
<td>8</td>
</tr>
<tr>
<td>SV, μl</td>
<td>9</td>
<td>8</td>
</tr>
<tr>
<td>TPR, mmHg/ml^-1/min</td>
<td>9</td>
<td>8</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Mean</th>
<th>CV, %</th>
<th>AD, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>MAP, mmHg</td>
<td>107±3</td>
<td>4.1±0.5</td>
<td>0.8±0.1</td>
</tr>
<tr>
<td>PI, ms</td>
<td>162±3</td>
<td>3.7±0.3</td>
<td>1.6±0.4*</td>
</tr>
<tr>
<td>CO, ml/min</td>
<td>76±2</td>
<td>7.2±0.7*</td>
<td>6.2±0.8*</td>
</tr>
<tr>
<td>SV, μl</td>
<td>203±5</td>
<td>7.0±0.5*</td>
<td>6.0±0.6*</td>
</tr>
<tr>
<td>TPR, mmHg/ml^-1/min</td>
<td>1.4±0.1</td>
<td>7.9±0.7*</td>
<td>6.0±0.5*</td>
</tr>
</tbody>
</table>

Values are means ±SE. Coefficients of variation (CV) and average beat-to-beat differences (AD) are shown as indexes of long-and short-term lability, respectively. MAP, mean arterial pressure; PI, pulse interval; CO, cardiac output; SV, stroke volume; TPR, total peripheral resistance; RBF, renal blood flow; MBF, mesenteric blood flow; HBF, hindquarter blood flow. * P < 0.05, significantly different from values of MAP in each group. † P < 0.95, significantly different between groups I and II.

Frequency bands are not complementary, and therefore spectral density power does not add to 100%.

To achieve a better resolution in the LF range, we also calculated spectra over 2,048 data points (~400 s). In some of these time series, power spectra of RBF showed a clear peak in the VLF band. No such VLF peaks were found in the power spectra of MBF or HBF. In general, the stationarity of these 400-s series was too low. Therefore the present frequency analysis was performed on time series of 512 data points (~100 s).

**Statistical analysis**

Data are presented as means ± SE except where noted. Statistical differences between coefficients of variation and average beat-to-beat differences of the hemodynamic parameters were assessed by a Wilcoxon nonparametric rank test. Analysis of variance was used to compare the results obtained by the spectral analyses for arterial pressure with those obtained for blood flows in the four defined frequency bands. In case of significance, a Bonferroni post hoc t-test was applied to identify the individual differences. Statistical significance was accepted at P < 0.05.
Results

Table 2.1 summarizes the 30-min average values and lability indexes of both the central and regional hemodynamic parameters. The steady-state values of arterial pressure were not different between groups I and II. However, average PI was slightly but significantly (-11%) lower in the group equipped for recording of ascending aortic flow. The reason for this discrepancy is not clear. Because we excluded fever, enhanced behavioral activity, or differences in respiration frequency as a possible cause (see METHODS, Protocol), the difference in heart rate is possibly related to the presence of the electromagnetic flow probe in the thoracic cavity. However, this difference in steady state between groups is of minor importance, since the lability indexes in Table 2.1 and dynamic fluctuations in PI (Table 2.2) were very similar. The long-term lability index of CO and the three regional blood flows exceeded those calculated for arterial pressure (P < 0.05 only for HBF and CO). Also the short-term lability indexes of regional blood flow, and of CO in particular, were significantly (P < 0.05) elevated above those obtained for arterial pressure.

Series of 512 beats of central and regional hemodynamic recordings as well as their respective power spectra are depicted in Figs. 2.1 and 2.2. These tracings demonstrate the different oscillatory patterns in the various hemodynamic signals. Although the rats were not moving, other events (like deep breaths) cause nonstationarities as indicated by the arrow in Fig. 2.2. Despite the increment in LF power caused by such an

<table>
<thead>
<tr>
<th>Table 2.2. Comparison of relative spectral powers in VLF, LF, MF, and HF bands for hemodynamic parameters in groups I and II</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter</td>
</tr>
<tr>
<td>-----------</td>
</tr>
<tr>
<td>n</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>MAP</td>
</tr>
<tr>
<td>PI</td>
</tr>
<tr>
<td>CO</td>
</tr>
<tr>
<td>SV</td>
</tr>
<tr>
<td>TPR</td>
</tr>
<tr>
<td>MAP</td>
</tr>
<tr>
<td>PI</td>
</tr>
<tr>
<td>RBF</td>
</tr>
<tr>
<td>MBF</td>
</tr>
<tr>
<td>HBF</td>
</tr>
</tbody>
</table>

Values are means ±SE. Frequency bands: VLF, very low; LF, low; MF, mid; HF, high. * P < 0.05, significantly different from values of MAP within a single frequency band within each group. † P < 0.05, significantly different between groups I and II.
event, most characteristic fluctuations were preserved. For each rat, mean power spectra were calculated by averaging the set of sequential spectra of the recording session. The resultant spectra were again averaged for all rats and are presented in Fig. 2.3. Figures 2.1 and 2.3 demonstrate the prevalence of LF and MF oscillations in the MAP signal and the absence of those oscillations in SV and in CO. Figures 2.2 and 2.3 illustrate the predominance of the LF fluctuations in MBF and to a lesser extent in RBF but not HBF. The average data for the different frequency bands are quantified in Table 2.2. The distribution of MAP and PI over the four different frequency bands was, except for MF of MAP, not different in both groups of rats, suggesting reproducible and comparable hemodynamic stability in the two sets of experiments. The magnitude of the respiratory (HF) fluctuations in CO was comparable to those in SV and made up approximately one-third of all variability in these signals. The HF oscillations in the peripheral flow signals were considerably smaller (~9%) and contributed little (~2%) to total MAP variability. MF variations were lowest for PI (~3%), and values ranging from 6 to 13% were found for the MF fluctuations in other parameters. LF fluctuations comprised ~35% of all variability in MAP. Because SV and CO displayed little LF spectral density power (~7%), the LF oscillations must have arisen from resistance to flow. LF oscillations were most pronounced in MBF (~35%), smaller in RBF and HBF (~17%), and lowest in SV and CO (~7%). VLF power was most abundant in the PI signal. High VLF values were also found for HBF but not for RBF, MBF, CO, and SV.

The vascular admittance gain of MAP and CO is presented in Fig. 2.4. In the HF band, the admittance gain of MAP vs. CO reached values up to 20, corroborating the impact of the respiratory fluctuations in CO on MAP in this frequency range. The
cross-correlation function between MAP and CO is presented in Fig. 2.5. Remarkably, fluctuations in MAP and CO behave independently and were only significantly correlated at zero delay, demonstrating the direct increase in arterial pressure as CO rises. The cross-correlation function between SV and PI is also depicted in Fig. 2.5. The positive correlation values about zero illustrate the lengthening of PI when SV increases and, vice versa, the increase of SV as PI gets large. In the peripheral vascular beds the influence of the respiratory cycle on blood flow variations gradually declined, as indicated by the decreasing values of the admittance gain at this frequency (Table 2.3).

The admittance gain of regional blood flow and MAP interactions is presented in Fig. 2.6. In most regional vascular beds, the fractional variations in blood flow in the MF and LF band exceeded those in arterial pressure (admittance gain > 1). Only for the LF band of the MAP-RBF interactions was the admittance gain smaller, but not significantly different from 1. Notably, the phase lag between arterial pressure and RBF and MBF in the LF band on the one hand was opposite to the phase shift between arterial pressure and HBF on the other (Table 2.3). This divergent behavior of the regional flow signals is further demonstrated in Fig. 2.7, showing the cross correlation with MAP. Both RBF and MBF are positively correlated at a time delay near zero, indicating that
changes in pressure are paralleled by similar changes in blood flow. In addition, significant negative correlations were found at approximately -2 s, meaning that 2 s after a pressure change, flow changes occurred in the opposite direction. In contrast, the average correlation with HBF at zero delay was negative, showing that in this vascular bed concurrent changes in pressure and flow occur in opposite directions.

**Discussion**

Beat-to-beat changes in arterial pressure, CO, and regional blood flows were analyzed by spectral analysis techniques to characterize the hemodynamics underlying systemic cardiovascular oscillations. The two main findings of this study are that 1) in conscious resting rats, the principal oscillations in systemic arterial pressure are centered about 0.13 Hz and do not arise from fluctuations in CO but, rather, from those in the renal and mesenteric vascular beds and 2) there is no evidence for autoregulation across all frequencies in any of the peripheral vascular beds studied.

In the present study in conscious rats, we used spontaneous variations of blood pressure as input for calculation of the transfer function to both CO as well as blood flow in three different vascular beds. Because coherence reached statistical significance for all frequencies in all vascular beds, it appeared that the naturally occurring fluctuations in blood pressure are adequate perturbations to study the transfer properties to the peripheral organs. In most previous related studies, interest was restricted to the analysis of pressure-flow interactions in the renal vascular bed [20, 21, 24, 25, 28-31], whereas transfer properties of arterial blood pressure to RBF were assessed in anesthetized
### Table 2.3. Average coherence, admittance gain, and phase shift for MAP vs. CO, RBF, MFB, and HFB

<table>
<thead>
<tr>
<th>Freq Band</th>
<th>Coherence</th>
<th>Admittance Gain, %</th>
<th>Phase, s</th>
</tr>
</thead>
<tbody>
<tr>
<td>MAP-CO</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>VLF</td>
<td>0.65 ±0.02</td>
<td>2.29 ±0.41</td>
<td>+0.03 ±0.48(^{a, c, e, d})</td>
</tr>
<tr>
<td>LF</td>
<td>0.65 ±0.02</td>
<td>1.84 ±0.30</td>
<td>+0.28 ±0.15(^{b, c, d})</td>
</tr>
<tr>
<td>MF</td>
<td>0.63 ±0.02(^{b})</td>
<td>3.13 ±0.27(^{b})</td>
<td>+0.003 ±0.03</td>
</tr>
<tr>
<td>HF</td>
<td>0.69 ±0.03</td>
<td>18.4 ±4.3(^{b, c, d})</td>
<td>-0.002 ±0.003(^{c, d})</td>
</tr>
<tr>
<td>MAP-RBF</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>VLF</td>
<td>0.64 ±0.04</td>
<td>1.06 ±0.17</td>
<td>+1.20 ±0.22</td>
</tr>
<tr>
<td>LF</td>
<td>0.76 ±0.04</td>
<td>0.91 ±0.10(^{c, d})</td>
<td>+0.96 ±0.11(^{c, d})</td>
</tr>
<tr>
<td>MF</td>
<td>0.79 ±0.03(^{a, d})</td>
<td>1.53 ±0.22(^{a, d})</td>
<td>+0.12 ±0.02(^{c, d})</td>
</tr>
<tr>
<td>HF</td>
<td>0.67 ±0.03</td>
<td>3.73 ±0.42(^{a})</td>
<td>+0.019 ±0.005</td>
</tr>
<tr>
<td>MAP-MBF</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>VLF</td>
<td>0.61 ±0.05</td>
<td>1.37 ±0.12</td>
<td>+1.67 ±0.22(^{a})</td>
</tr>
<tr>
<td>LF</td>
<td>0.76 ±0.04(^{b})</td>
<td>2.20 ±0.24(^{b})</td>
<td>+0.90 ±0.10(^{c, d})</td>
</tr>
<tr>
<td>MF</td>
<td>0.74 ±0.04(^{b})</td>
<td>1.84 ±0.14</td>
<td>-0.10 ±0.04(^{b})</td>
</tr>
<tr>
<td>HF</td>
<td>0.68 ±0.05</td>
<td>4.46 ±0.57(^{a})</td>
<td>+0.025 ±0.006(^{a})</td>
</tr>
<tr>
<td>MAP-HBF</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>VLF</td>
<td>0.63 ±0.05</td>
<td>3.44 ±0.56</td>
<td>+0.61 ±0.39</td>
</tr>
<tr>
<td>LF</td>
<td>0.66 ±0.06</td>
<td>2.30 ±0.26(^{b})</td>
<td>-0.50 ±0.16(^{a, b, c})</td>
</tr>
<tr>
<td>MF</td>
<td>0.62 ±0.05</td>
<td>3.15 ±0.52(^{b})</td>
<td>-0.09 ±0.03(^{b})</td>
</tr>
<tr>
<td>HF</td>
<td>0.67 ±0.05</td>
<td>6.87 ±0.58(^{a})</td>
<td>+0.037 ±0.007(^{a})</td>
</tr>
</tbody>
</table>

Values are means ± SE. \(^{a, b, c, d}\)Significantly different from values within the same frequency band of MAP-CO, MAP-RBF, MAP-MBF, and MAP-HBF, respectively.

Animals during forcing of the arterial pressure signal\(^{24, 25, 30}\).

As indicated by the individual tracings in Figs. 2.1 and 2.2, three distinct oscillations in MAP can be identified under resting conditions in rats. The hemodynamic characterization of these HF, MF, and LF oscillations in MAP is discussed in detail below. For reasons explained in METHODS VLF fluctuations are not discussed.

**Respiratory (HF) oscillations in MAP**

The amount of the respiratory-related HF power to overall MAP variability was low and generally not higher than 5%, which is roughly similar to values found in other rat studies\(^{9, 11, 12, 14}\). Particularly when longer time tracings were analyzed, the contribution of HF power to total MAP variability declined because in resting rats the breathing rhythm was regularly interrupted, about every minute, by a deep breath, causing nonstationarities in the hemodynamic recordings as indicated in Fig. 2.2. The present data clearly indicate that changes in CO were the primary source of HF fluctuations in MAP. Comparison of the vascular admittance values in the HF band indicated to what degree these respiratory fluctuations in CO were dampened along the vascular tree. The HF fluctuations in HBF, MFB, and RBF were respectively 2.7, 4.1, and 4.9 times smaller than those in the ascending aortic flow. Regarding the similarity in the tracings of SV and CO (Fig. 2.1) and the difference in HF spectral power between SV and PI, it can
be concluded that, at least in resting conditions, the HF fluctuations in CO were mainly caused by respiration-synchronous oscillations in SV and not in PI. Because neither sinoaortic denervation nor chronic sympathectomy reduced the magnitude of the HF oscillations in MAP in rats\(^9\), it seems that in this species the respiratory-related fluctuations in CO are devoid of neural influences and are purely mechanical in origin. In fact, as indicated by the positive cross-correlation function, the lengthening of the interbeat interval may have a stabilizing influence on the variability in CO when SV rises\(^9\). In dogs, however, heart rate oscillations seem to dominate the lability in CO\(^7\), whereas in humans, as in rats, fluctuations in SV are the main source of respiratory-related variability in CO\(^9\).

**Autonomic (MF) oscillations in MAP**

In rats, MF oscillations have been shown to be largely dependent on the integrity of the autonomic nervous system\(^9\). In the present study spectral density power and vascular admittance of the MF band were roughly similar for ascending aortic and regional blood flows. This suggests that both MF fluctuations in CO as well as resistance to flow contribute to these ~ 2.5 s oscillations in MAP. The extent that MF power contributed to total MAP variability was ~ 10%. This seems rather small compared with other rat studies\(^1\), but can be explained by the differences in the definition of the width of this frequency band in these studies as well as the arousal level of the animals\(^1\). In sympathectomized rats, MF oscillations in MAP were absent\(^1\), suggesting that these MF oscillations depend highly on the integrity of the sympathetic nerves. However, in direct recordings of sympathetic nerve activity in rats, no explicit peaks at 0.4 Hz were found\(^4\). Further experiments are needed to clarify whether this discrepancy could be explained by transfer properties of the nerve signal to vascular smooth muscle or to cardiac muscle. Vascular MF admittance gain was higher in the hindquarter than renal or mesenteric vascular beds, which may be indicative of a greater autonomic influence on this vascular bed in the resting state\(^5\). This hypothesis is supported by the present findings. As indicated by the cross-correlation function, HBF and MAP were significantly negatively correlated about zero time delay. A physiological interpretation of this relationship is that the HBF changes are responsible for the MAP changes, which may be governed by the autonomic nervous system. In contrast, RBF and MBF showed significant positive correlations with MAP about zero. This probably reflects the elastic behavior of these vascular beds when pressure rises. The above-mentioned differential pattern of pressure-flow interactions in the three vascular beds only applies to the resting state and may be different when behavior changes. We found that the occasional movements of the rats increased blood flow through this region by > 40%, whereas RBF and MBF fell only by ~ 3 and 9%, respectively. This may explain the significantly higher long-term lability index for HBF. Furthermore, it should be noted that blood flow through the abdominal aorta is the result of flow through the hindlimb muscles, tail, and parts of the skin.

**Myogenic (LF) oscillations in MAP**

This is the first study in rats showing that a substantial part of LF power in MAP was due to clear oscillations centered about 0.13 Hz. Spectral analysis of the hemodynamic recordings indicated that the LF oscillations in MAP resulted from resistance to flow and were especially prominent in the mesenteric vascular bed. Because
this vascular bed receives a relatively high proportion of CO in resting rats, it is not surprising that these LF oscillations in MBF were transmitted into the MAP signal. LF fluctuations in RBF may have also contributed, since the time lags of the LF oscillations in RBF and MBF with MAP were very similar. The time shift between MAP and the regional flow signals identified by the cross-correlation function were somewhat different, because this calculation necessarily includes all frequency components. Nonetheless, we found that the fluctuations in MAP were leading those in MBF and RBF by ~2 s and that they were negatively correlated. This correlation pattern was repeated every 8 s, as could be expected from the predominance of the LF fluctuations in these signals. The physiological interpretation of such a pressure-flow relationship is that when pressure increases, significant reductions in blood flow occur at the indicated delay. The present characteristics are suggestive of active myogenic control of blood flow. In fact, in this conscious preparation, the myogenic response of the mesenteric arteries seems so robust that the LF fluctuations in MBF were proportionally about twice the MAP fluctuations. Thus rather than autoregulation, this overshooting reaction points to superregulation. The pressure-flow interactions seem less vigorous in the renal vascular bed, since here the admittance gain was smaller but not statistically significantly different from 1. In anesthetized rats, with the use of forced pressure fluctuations, similar interactions between arterial pressure and RBF have been identified in the 0.1- to 0.2-Hz band with a response time of ~1 s. In addition, in various experimental and clinical microcirculatory preparations (muscle, cerebral, and skin), blood flow shows spontaneous fluctuations in frequency ranges similar to LF oscillations seen here. Interestingly, these fluctuations increased in frequency and amplitude under conditions of low blood pressure or hypoperfusion, suggesting a myogenic reaction.

The LF oscillations in regional blood flow are not necessarily pressure dependent but may also arise from an endogenous oscillator located in the peripheral vessels. Resistance arteries isolated from various organs, in particular from the mesentery, are known to exhibit spontaneous LF oscillatory contractions, especially when stimulated with agonists. In addition, in a modeling study, Ursino and colleagues showed that self-sustained oscillations can develop when the dynamic component of the myogenic response of terminal arterioles is greater than that of more proximal arterioles in a vascular bed.

Because the present LF oscillations originate from resistance vessels, their activity must be somewhat synchronized or otherwise no rhythmicity would have been found in whole organ blood flow. Thus the absence of peaks in this frequency band may not decisively exclude myogenic activity, because the contractile activity of the smooth muscle cells may be out of phase. However, from microcirculatory work it is known that periodic vasomotion may spread upstream over the arteriolar network. This property may synchronize and increase the amplitude of the oscillation, so that it can be detected in whole organ blood flow. Similarly, related oscillations in tubular pressure in adjacent nephrons are known to entrain and thereby enlarge their impact on autoregulation of RBF.

The superregulatory behavior of the LF oscillations does not exclude its serving a homeostatic function in the control of organ blood flow. Although its characteristics are that of a vicious cycle, causing a further rise in resistance in response to a pressure
increase, this mechanism is probably often overruled by the baroreflex when systemic pressure increases its threshold value. On the other hand, in the case of a pressure decrease, a further pressure fall is probably also prevented by the baroreflex. Support for the stabilizing function of the autonomic nervous system comes from a study in sympathectomized rats. Arterial pressure lability in these rats is characterized by many spontaneous hypotensive episodes. The regional flow pattern accompanying such a hypotensive event showed that vasodilation in the mesenteric vascular bed followed the pressure fall amplifying the fall in pressure.

The amplitude of the LF oscillations may also be influenced by other oscillatory mechanisms. Yip and colleagues reported that during inhibition of slow (~0.03 Hz) tubuloglomerular feedback (TGF) related oscillations in single-nephron blood flow fluctuations in the LF myogenic band increased, whereas they fell at saturation of TGF. On the basis of such an inverse relationship, it is tempting to speculate that the abundance of LF power in the mesenteric vascular bed may be partly due to the absence of a slow regulatory mechanism in this organ.

In humans, the so-called 10-s rhythm in blood pressure and heart rate has been thought to depend primarily on the autonomic nervous system activity. Changes in the amplitude of these LF oscillations are increasingly used as markers to estimate clinical autonomic disturbances. The clinical implication of this study is that if the elements of the arterial tree in humans behave similarly as those in rats, myogenic fluctuations may also contribute to the so-called 10-s rhythm in blood pressure in humans.

In summary, in conscious resting rats, spontaneous fluctuations in systemic arterial pressure predominantly exhibit slow (~8 s) oscillations, which do not arise from fluctuations in CO but originate from regionally specific oscillatory mechanisms contributing to resistance to flow. These blood flow fluctuations were most pronounced in the mesenteric and renal vascular beds and displayed a flow-conserving myogenic-like superregulatory character. In contrast, blood flow oscillations in the hindquarter vascular bed were typified as being caused by mechanisms that serve systemic pressure control.

References


Chapter 2

Chapter 3
Validation of a continuous baroreceptor reflex sensitivity index calculated from spontaneous fluctuations of blood pressure and pulse interval in rats

Jan Oosting
Harry A.J. Struijker-Boudier
Ben J.A. Janssen

Published in: Journal of Hypertension 1997, 15:391-399
Chapter 3

Abstract

Objective To develop and validate a technique for the continuous computerized calculation of the baroreceptor reflex sensitivity (BRS) of the heart rate in rats.

Design The BRS was calculated from spontaneous changes in blood pressure and pulse interval using spectral analysis as well as time-series techniques. The BRS values obtained with these techniques were compared with those obtained by standard pharmacological methods.

Methods The blood pressure and pulse interval in adult Wistar-Kyoto (WKY) rats were recorded on a beat-to-beat basis for two consecutive 30-min periods. During one of these periods the BRS was determined pharmacologically by injections of nitroprusside and phenylephrine. Measurements were performed after administration of saline as vehicle or during manipulation of the autonomic nervous system by infusion of metoprolol, methyl-atropine and hexamethonium. Sequential time-series methods for continuous BRS calculation were tested for 24 h periods in intact WKY rats as well as in WKY rats that had been subjected to sino-aortic denervation or to electrical lesioning of the nucleus tractus solitarius.

Results The correlation coefficient between BRS values in intact WKY rats derived from the pharmacological method and those from spectral analysis techniques was low ($R^2 = 0.16$). The correlation coefficient between BRS values from the pharmacological method and those from the developed time-series method was higher ($R^2 = 0.64$). The BRS measured using the latter method was found to vary over 24 h with the highest values during the sleeping period. After surgical elimination of the baroreflex, the algorithm returned BRS values close to zero throughout the 24 h period. The BRS estimate was found to be a measure of the parasympathetic rather than of the sympathetic component of the baroreceptor reflex.

Conclusion The developed time-series method calculates an index of the gain of the cardiac baroreflex in rats faithfully. This method can be implemented in data acquisition software, allowing continuous on-line monitoring of the cardiac baroreflex gain.
Validation of continuous BRS measurements

Introduction

The baroreceptor reflex is the most important regulatory mechanism in the short-term control of the heart rate and blood pressure and operates through the autonomic nervous system. In the past, the gain of the cardiac baroreflex, further referred to as the baroreflex sensitivity (BRS), has usually been obtained by comparing changes in heart rate as a consequence of imposed blood pressure changes induced by pharmacological or mechanical methods. Measuring the BRS from spontaneous variations in blood pressure and heart rate has, in principle, several advantages over methods that provoke changes in blood pressure. There is no need to administer vasoactive compounds or external appliances that could influence the baroreceptor reflex by a direct action on receptor or effector sites. Second, the BRS is measured within physiological blood pressure ranges, allowing the computation of the gain at the operating point of the blood pressure with minimal non-specific effects from other afferent nerves. Furthermore, these methods do not arouse subjects or animals, thereby reducing stress-induced effects. Finally, in contrast to the pharmacological or mechanical methods, they are suitable to assess the BRS over prolonged periods of time.

Published methods that evaluate the BRS from spontaneous changes in blood pressure and heart rate make use of spectral analysis techniques, of spontaneously occurring ramps in blood pressure or of statistical relationships between blood pressure and pulse interval changes. With advancing computer technology, it should now be possible to implement such methods in data-acquisition software, making continuous BRS recordings possible. However, before such a method can be applied, not only technical but also physiological testing is required. Here we describe such physiological validation studies. Experiments were performed in rats. This species was chosen to allow extensive surgical and pharmacological interventions to manipulate the gain of the baroreflex. We compared spectral analysis techniques with standard pharmacological methods in terms of their applicability for the on-line determination of the cardiac baroreflex gain. In addition we evaluated the effect of low-pass filtering the spontaneous fluctuations of the blood pressure and heart rate data. There is increasing evidence that the baroreflex is especially capable of buffering low-frequency variations in pressure. After sino-aortic denervation (SAD) the spectral density power of the blood pressure increased, especially at low frequencies (below 0.1 Hz), whereas it decreased at high frequencies.

The BRS calculated from spontaneous fluctuations should show a high correlation to pharmacological methods, detect contrasts in animals with known differences in BRS and return a zero value in cases without baroreflex activity. For these reasons the methods were tested on data obtained in rats that had been treated with agents that block part or all of the autonomic nervous system as well as in rats that had been subjected to sino-aortic denervation or to bilateral lesioning of the nucleus tractus solitarius. With regard to these criteria we developed an algorithm that calculated faithfully and continuously a minute-to-minute BRS index from spontaneous ramps in blood pressure and pulse interval in 16 rats simultaneously using an IBM-compatible 50 MHz 486 computer.
Chapter 3

Methods

Animals

Adult WKY rats weighing 280-300 g each were used. The rats were obtained from the central Animal Facilities of the Universiteit Maastricht. Experiments were performed according to the Universiteit Maastricht guidelines. After they had undergone surgery, the animals were housed individually in cages. Rats were kept on a 12 h/12 h light/dark cycle and were allowed normal rat chow and drinking water ad libitum.

Surgery

All rats were instrumented with an arterial catheter for measuring their blood pressure and two venous catheters for administering drugs, as described in detail previously. In short, the catheters were implanted with the rats under pentobarbital anaesthesia (60 mg/kg intraperitoneally) via the femoral vessels and then tunneled to the lower back of the animal. The catheters were exteriorized through a 40 cm long steel spring that was fixed to the back muscles using silicone rubber attached to a piece of gauze. The venous catheters were filled with 5 U/ml heparinized saline solution. The steel spring was led to the outside of the cage, where the arterial catheter was connected via a hydraulic swivel (model 375/20; Instech Labs, Plymouth Meeting, Pennsylvania, USA) to a low-volume displacement pressure transducer (microswitch, model 156PC 156WL; Honeywell, Inc., Amsterdam, The Netherlands). The arterial canula was kept patent by continuous arterial infusion of 30 U/ml heparinized saline solution at a rate of 2.4 ml/day. This set-up allowed us to measure blood pressures continuously and to administer drugs without disturbing the rats.

Data acquisition

After at least 3 days of recovery the measurements were performed in the home cage of the animal. The arterial pressure transducer was connected to an amplifier that delivered a high-voltage signal to an analogue-to-digital converter board (model 2814; Data Translation, CN Rood, Rijswijk, The Netherlands) connected to an IBM-compatible computer. The blood pressure signal was sampled with two computerized units, one operating at 500 Hz, the other at 1000 Hz. The detection limit of blood pressure changes was smaller than 0.1 mmHg. Beat-to-beat pulse interval, heart rate and mean arterial pressure values were calculated on-line and stored on hard disc. We recorded the mean arterial pressure instead of the systolic blood pressure, because, especially during long-term recordings, the mean arterial pressure is more reliable than is the systolic blood pressure. The pulse pressure in unobserved catheterized rats sometimes decreases due to movement artefacts and catheter position problems. The mean arterial blood pressure is less affected by this than is the systolic blood pressure.

Protocols

To validate the methods which calculate an index of the gain of the cardiac baroreflex from spontaneous blood pressure fluctuations, a set of beat-to-beat blood pressure and pulse interval sequences was obtained for rats in which the baroreflex sensitivity was also measured by the pharmacological "Oxford method". In order to obtain a wide range in the cardiac baroreflex gain, recordings were performed under the
Validation of continuous BRS measurements

following conditions: the cardiac sympathetic system was blocked by 1.5 mg/kg metoprolol (n = 10); the cardiac parasympathetic system was blocked by 0.5 mg/kg methyl- atropine (n = 7); both cardiac nervous systems were blocked by combining metoprolol and methyl-atropine (n = 11); ganglionic transmission was blocked by hexamethonium (n = 14); and saline was administered as vehicle (n = 26). Metoprolol and methyl-atropine were dissolved in saline and injected as an intravenous bolus (<350 µl). To avoid great falls in blood pressure, which are evoked when hexamethonium is administered as a bolus injection, we infused this agent at a rate of 25 mg/kg per h, starting 1 h before the recordings. When rats were used for multiple recordings, the recordings of the separate autonomic conditions were performed at intervals of at least 2 days.

Under each condition two consecutive recordings each of duration 30 min were performed in random order. During one period the blood pressure and heart rate were allowed to vary spontaneously. During the other the blood pressure was changed by bolus injections of 0.1 mg/ml sodium nitroprusside and 0.03 mg/ml phenylephrine. These agents were injected alternately in volumes ranging from 2 to 30 µl, administered within 2-5 s to obtain changes in blood pressure ranging from -40 to +40 mmHg. At least six injections of each drug were administered. The injection volumes were allocated in random order.

In an additional experiment surgical disruption of the baroreflex was performed in six WKY rats 1 week before the catheters were implanted. Three rats were subjected to SAD using the method described by Krieger. A bilateral electrical lesion of the nucleus tractus solitarius (NTS) was made in 3 other rats according to the method of Buchholz and Nathan. That denervation had been achieved in these rats was confirmed by the absence of heart rate changes greater than 15 beats/min after a reduction by about 40 mmHg in blood pressure had been induced by injections of sodium nitroprusside. In addition, the presence of NTS lesions was confirmed by histology. In these rats as well as in intact WKY rats 24 h beat-to-beat recordings of blood pressure and heart rate were performed.

Calculations of the BRS of the heart rate

Pharmacologically induced blood pressure changes

From each change in pressure evoked by bolus injections of phenylephrine or sodium nitroprusside, the BRS was calculated as the slope of the linear regression function between the beat-to-beat changes in mean arterial pressure and the related beat-to-beat changes in pulse interval at a delay that scored the highest correlation coefficient. The average BRS estimate was calculated as the mean value of 8 ± 1 increasing and 11 ± 1 decreasing ramps. In intact rats, the gains of the increasing and decreasing ramps did not differ (phenylephrine 0.94 ± 0.11, sodium nitroprusside 0.95 ± 0.09). The delay of the pulse-interval response was in the range 2-14 beats, tachycardic responses generally being two or three beats later than bradycardic responses. Because methods triggering on spontaneous fluctuations provide only a value of the gain of the baroreflex, we did not construct a complete sigmoidal baroreflex curve to assess the full range of the heart-rate response.
Chapter 3

Spontaneous blood pressure changes

Initially the BRS was calculated from the spontaneous fluctuations in blood pressure by two methods. First we tested the spectral analysis technique described by Robbe et al., in which the transfer function between variations in blood pressure and in heart rate in the midfrequency band was calculated as an index of the BRS. We divided the time series into consecutive parts of 256 beats. These segments were subjected to cosine-tapering of 30 data points. Then the transfer function was calculated using the blood pressure as the input and the pulse interval as the output signal. The average pulse interval during the recording was taken as the sample frequency. Results were averaged over all consecutive data blocks and the mean value of the gain in the frequency range 0.35-0.65 Hz was used as an index of the BRS. In rats the variability in this frequency range is generally attributed to the baroreflex.

The first time-series method we tested was a direct translation of the original method described by Bertinieri et al. Their algorithm searches for periods of three or more consecutive beats in the blood pressure and pulse interval signal within which the blood pressure changes by at least 1 mmHg per beat, while the pulse interval changes in the same direction during the following beat by at least 4 ms. The slope of the linear correlation function of the two parameters in such a ramp is taken as an index of the BRS. This method has been applied successfully to data obtained from humans and cats. Our analysis on data obtained from rats, however, revealed that, with the original threshold pressure (1 mmHg) and pulse interval (4 ms) settings, it was impossible to find a reasonable number of ramps to allow continuous registration of the BRS and a reliable comparison with pharmacological BRS estimates. There are several physiological reasons for this. First of all, the average heart rate and respiration rate are much higher in rats than those in cats and humans. Thus, the algorithm of the time-series method should be adapted using appropriate delays between changes in pressure and in pulse interval. Using the beat-to-beat tracings during injections of phenylephrine and nitroprusside, we found that baroreflex-related changes in pulse interval became apparent 3-5 beats after a pressure change. This delay corresponds to a time shift roughly in the range 0.6-1.2 s. Second, because of the higher cardiac frequency, the beat-to-beat differences in blood pressure and heart rate are smaller than those in species with lower heart rates. We tried several threshold settings for the pressure and pulse interval changes to trigger a BRS calculation in combination with the 3-5 beats delay. However, it appeared that the height of the thresholds did not lead to the detection of longer ramps in these unfiltered data sets. Rather, we observed that the fast respiratory-related variations in blood pressure and pulse interval hampered the detection of ramps longer than five beats during which the blood pressure or pulse interval increased or decreased continuously (see Fig. 3.1). This was one of the reasons to apply a filtering procedure to the rat data set. The physiological validity of this filtering procedure has been supported by several recent observations. First, evidence is accumulating that the baroreflex buffers predominantly low-frequency (< 0.1 Hz) fluctuations in blood pressure. Following SAD in cats, rats and dogs, the spectral density power of low- rather than of high-frequency oscillations in arterial pressure increased. In rats, but not in cats, also the spectral power density of the high-frequency pulse-interval oscillations increased after SAD and the coherence between respiratory-related fluctuations in pressure and in pulse interval did not alter. This suggests that
Validation of continuous BRS measurements

![Graph showing PI (ms) and MAP (mmHg) over beats](image)

**Figure 3.1.** Beat-to-beat pulse interval (PI) and mean arterial pressure (MAP) values shown as they were recorded originally (thin line) and after the application of a 10-beat moving average function (thick line). The arrows indicate the first beat for which the algorithm found a ramp that was included into the calculation of the baroreflex sensitivity estimate.

Baroreflex buffering of the heart rate in the rat is not the major determinant of the high-frequency oscillations in heart rate. In support of this view, we found recently that respiratory-related fluctuations in pressure and cardiac output in rats were caused mainly by oscillations in stroke volume rather than by baroreflex-mediated changes in pulse interval. Also in humans, it has now been demonstrated that respiratory sinus arrhythmia contributes to such arterial pressure fluctuations, rather than that it represents baroreflex buffering. Thus, in the rat, the fluctuations in blood pressure and heart rate related to the respiratory cycle are probably due more to mechanical than to nervous mechanisms. On the basis of these arguments we chose to eliminate the relatively fast fluctuations from the blood pressure and pulse interval signal by low-pass filtering of the pressure and pulse interval signal using a 10-beat moving average function. Thus, we restricted the focus of our analysis to the low-frequency part of the spectrum in rats. The efficacy of this procedure is visualized in Figure 3.1. The result is that the blood pressure and pulse interval signals show 'longer lasting' transients, allowing the algorithm to find triggers.

As a last point, an intrinsic property of the available time-series methods is that they introduce bias in the sense that ramps in blood pressure and pulse interval are
Chapter 3

included in the calculation of the BRS only when they have a certain magnitude and direction. The magnitude of bias introduced by this is not known. We chose to circumvent this problem by including in the calculation all ramps in the pressure signal, neglecting the direction and magnitude of the change in pulse interval. The idea behind this approach is that, if baroreflex-mediated changes in pulse interval are present, then they should appear as such when the blood pressure changes. In all other cases the blood pressure and heart rate fluctuations were assumed to occur randomly, thus only adding noise to the detection system. The same assumption also applies when spectral analysis techniques are employed.

On the basis of the aforementioned criteria, we chose to implement the following algorithm. Apply a moving average of 10 beats both to the blood pressure signal and to the pulse interval signal. Search for ramps of decreasing or increasing blood pressure of four beats or more. When a ramp has been found, determine the slopes between the blood pressure ramp and pulse interval changes with three, four and five beats delay. Calculate the BRS as the average value of these three slopes.

By using this algorithm we found that the average ramp length in intact WKY rats was 9.7 ± 1.6 beats (mean ± SD), whereas the average blood pressure change was 3.4 ± 0.9 mmHg for ramps during which the blood pressure increased and 3.8 ± 0.8 mmHg for ramps during which the blood pressure decreased. On the average an increase in blood pressure was associated with a higher BRS estimate (0.32 ± 0.03 ms/mmHg) than was a decrease in blood pressure (BRS 0.17 ± 0.03 ms/mmHg). Of all beats, 49.8 ± 4.1% were included in the calculations. Because the BRS is calculated as the regression coefficient of the pulse interval and pressure changes over several beats in a ramp, the BRS estimate can be lower than 1 ms/mmHg, despite the 2 ms sampling rate.

One of the technical considerations for adapting the timeseries method rather than the spectral analysis method was that we wanted to calculate the BRS on-line. The calculations of the BRS index can be performed as soon as a blood pressure ramp has been found, imposing only a small time penalty. Frequency analysis requires stationary signals with a certain minimum length to detect the frequencies of interest. Therefore, these latter methods can be applied only to certain time periods and take relatively more time. In general, the algorithm detected about 20 ramps per minute in the blood pressure time series.

Drugs

Sodium pentobarbital was obtained from Sanofi Sante BV (Maassluis, The Netherlands). Metoprolol tartrate (metoprolol), aropine methyl nitrate (methyl-atropine), hexamethonium bromide (hexamethonium) and phenylephrine hydrochloride (phenylephrine) were obtained from Sigma Chemical Co. (St Louis, Missouri, USA). Sodium nitroprusside (nitroprusside) was obtained from Janssen Chimica (Beerse, Belgium).

Statistics

Differences between baseline blood pressure and heart rate values obtained during the two consecutive 20 min periods were compared using a paired Student's t-test. BRS values obtained from the methods using spontaneous fluctuations in blood pressure and pulse interval were compared with those from the pharmacological method by Pearson correlation. The correlation, slope and intercept were computed. The magnitudes
Validation of continuous BRS measurements

Table 3.1. Baseline haemodynamic values in groups of rats treated with autonomic blocking agents

<table>
<thead>
<tr>
<th></th>
<th>Arterial pressure (mmHg)</th>
<th>Heart rate (beats/min)</th>
<th>Baroreflex sensitivity (ms/mmHg)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>control</td>
<td>drug</td>
</tr>
<tr>
<td>Metoprolol</td>
<td>10</td>
<td>108±3.9</td>
<td>109±4.1</td>
</tr>
<tr>
<td>Methyl-atropine</td>
<td>7</td>
<td>110±4.6</td>
<td>115±2.5</td>
</tr>
<tr>
<td>Metoprolol + Methyl-atropine</td>
<td>11</td>
<td>119±4.8</td>
<td>128±4.5</td>
</tr>
<tr>
<td>Hexamethonium</td>
<td>14</td>
<td>144±4.0</td>
<td>115±2.9*</td>
</tr>
</tbody>
</table>

Baroreflex sensitivity was assessed by a pharmacological method (see text) Values are expressed as means ± SEM. * P < 0.05, versus control by paired Student’s t-test

of the differences between groups were compared using analysis of variance. P < 0.05 was considered statistically significant.

Results

Under all treatments baseline blood pressure and heart rate values did not differ between the two 30 min recording periods (P > 0.6). Therefore pooled data are presented in Table 3.1. In addition the average BRS values calculated by the pharmacological method are shown. The baseline mean arterial pressure values varied between the groups of rats. However, under control conditions, average heart rates and average BRS values were similar in the groups. As expected, hexamethonium administration reduced the arterial pressure, whereas metoprolol decreased and methyl-atropine increased the heart rate. The BRS fell slightly during metoprolol administration, but was suppressed severely during administration of methyl-atropine, of the combination of methyl-atropine and metoprolol and of hexamethonium.

The linear correlation between BRS values calculated by pharmacological means and by the spectral analysis method is presented in Figure 3.2. Figure 3.2 shows that the spectral method yielded variable gains, even under conditions in which the baroreflex was blocked. The correlation coefficient was low (R² = 0.16) and the intercept was at 0.66 ms/mmHg (Table 3.2, experiment 1). The linear correlation between BRS values calculated by pharmacological means and the time-series method using nonfiltered blood pressure and heart rate signals was not much better than that obtained with the spectral method. As indicated (Table 3.2, experiments 2 and 3), the correlation coefficient between the BRS estimates increased from R² = 0.09 to R² = 0.41 when, instead of a one-beat delay, the average of the BRS estimates calculated with three, four and five beats delay between the mean arterial pressure and the pulse interval was used. The correlation coefficient was not further increased by setting specific thresholds for the mean arterial pressure and pulse interval (data not shown).

Using the arguments indicated in the methods section, we applied moving-average filters to the mean arterial pressure and pulse interval signals and examined whether this would improve the correlation with the pharmacological BRS estimate. We found
Figure 3.2. Comparison between baroreflex sensitivity of the heart rate (BRS) values calculated by pharmacological methods and by spectral analysis techniques (using a transfer function). The methods were compared during control (Circle) and treatment with metoprolol (Square), methyldigoxine (Upward triangle), and a combination of metoprolol and methyldigoxine (Downward triangle). The drawn line represents the regression function (correlation $R^2=0.16$, constant $0.66 \pm 0.84$, slope $0.98 \pm 0.42$)

Figure 3.3. Comparison between baroreflex sensitivity of the heart rate (BRS) values calculated by pharmacological methods and the adapted time-series method (method 7, Table 2). Symbols represent the same conditions as in figure 3.2. The drawn line represents the regression function (correlation $R^2=0.64$, constant -0.06 $\pm 0.19$, slope 0.35 $\pm 0.03$)

that, with a 10-beat moving-average procedure, the correlation between the BRS estimates rose to $R^2=0.64$ (Table 3.2, experiment 7). We tested also different lengths of the moving average procedure (5 or 12 beats, instead of 10 beats, see Table 3.2, experiments 5 and 6). Such differences in smoothing had only minor effects on the correlation between the BRS estimates. To investigate whether the algorithm could be further improved, several other settings of variables in $i$ were tested. When the delay was made variable in the range 3-5 beats and the linear relation between the blood pressure and the pulse interval was calculated at the delay with the highest correlation, the correlation to the pharmacological method fell slightly to $R^2=0.46$ (Table 3.2, experiment 7d). However, in this case also SAD and NTS-lesioned animals exhibited a positive BRS (0.37 $\pm 0.06$ ms/mmHg). When ramps were used in which the blood pressure change was greater than 2 mmHg, then the correlation coefficient was lowered slightly to $R^2=0.55$ (Table 3.2, experiment 7a) and the number of detected ramps fell from 17.7 $\pm 3.2$ to 8.9 $\pm 2.4$/min (mean $\pm$ SD). When the pulse interval was set to change by more than 1 ms in the same direction as the blood pressure ramp, not only did the correlation decrease but also the intercept increased (Table 3.2, experiment 7b). Finally, we selected ramps in which the correlation function between the blood pressure and the pulse interval ramp
Table 3.2. Comparison of the pharmacological method and techniques calculating the baroreflex sensitivity of the heart rate from spontaneous changes in arterial pressure and pulse interval.

<table>
<thead>
<tr>
<th>Exp</th>
<th>Description</th>
<th>$R^2$</th>
<th>Constant</th>
<th>Slope</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Transfer function analysis [6]</td>
<td>0.16</td>
<td>0.66±0.84</td>
<td>6.98±0.42</td>
</tr>
<tr>
<td>2</td>
<td>No smoothing, one beat delay</td>
<td>0.1</td>
<td>0.00±0.23</td>
<td>0.18±0.05</td>
</tr>
<tr>
<td>3</td>
<td>No smoothing, 3-5 beats delay</td>
<td>0.41</td>
<td>-0.04±0.10</td>
<td>0.21±0.03</td>
</tr>
<tr>
<td>4</td>
<td>Moving average 10 beats, one beat delay</td>
<td>0.14</td>
<td>-0.05±0.17</td>
<td>0.18±0.05</td>
</tr>
<tr>
<td>5</td>
<td>Moving average 5 beats, 3-5 beats delay</td>
<td>0.61</td>
<td>-0.08±0.14</td>
<td>0.34±0.03</td>
</tr>
<tr>
<td>6</td>
<td>Moving average 12 beats, 3-5 beats delay</td>
<td>0.52</td>
<td>-0.08±0.11</td>
<td>0.31±0.03</td>
</tr>
<tr>
<td>7</td>
<td>Moving average 10 beats, 3-5 beats delay</td>
<td>0.64</td>
<td>-0.06±0.10</td>
<td>0.35±0.03</td>
</tr>
<tr>
<td>7a</td>
<td>Change in blood pressure within ramp &gt; 2 mmHg</td>
<td>0.55</td>
<td>-0.06±0.12</td>
<td>0.33±0.03</td>
</tr>
<tr>
<td>7b</td>
<td>Change in pulse interval within ramp &gt; 1 ms</td>
<td>0.48</td>
<td>0.38±0.30</td>
<td>0.74±0.09</td>
</tr>
<tr>
<td>7c</td>
<td>Correlation between pulse interval and blood pressure within ramp &gt; 0.5</td>
<td>0.79</td>
<td>0.42±0.35</td>
<td>0.72±0.11</td>
</tr>
<tr>
<td>7d</td>
<td>Best linear fit between 3 and 5 beats delay</td>
<td>0.46</td>
<td>0.45±0.06</td>
<td>0.10±0.19</td>
</tr>
</tbody>
</table>

was higher than $R^2 = 0.25$. This method also gave a lower correlation to the pharmacological method and a high constant of the correlation function (Table 3.2, experiment 7c). For these reasons we finally chose to implement the algorithm of experiment 7 (Table 3.2, Fig. 3.3).

To illustrate the efficacy of this algorithm and its on-line implementation, we compared the 24 h BRS values for WKY rats, SAD and NTS-lesioned rats (Fig. 3.4). Figure 3.4 shows that the BRS in intact WKY rats was higher during the light period than it was during the dark period and that the BRS in SAD and NTS-lesioned animals was close to zero throughout the 24 h.

Discussion

In this study we have validated a sequence method that can be used for the continuous calculation of an index of the baroreceptor reflex activity from spontaneous fluctuations in blood pressure and pulse interval in rats. The method yields predominantly estimates of the gain of the parasympathetic component of the cardiac baroreflex which correlate well with those obtained by pharmacological means. The algorithm can be implemented online in data-acquisition software.

Our study indicates that, in rats, in contrast to in humans, BRS values obtained by calculation of the transfer function between blood pressure and pulse interval oscillations in the mid-frequency bands were correlated poorly to those acquired by pharmacological techniques. This could be a species-specific problem and might depend on the frequency range within which the baroreceptor reflex in rats is active. Cerutti et al. have shown that approximately 50% of the oscillations in the range 0.27-0.74 Hz can be attributed to the baroreflex. Our data indicated that, although the absolute values of the reflex gains in intact rats were comparable between the pharmacological and spectral methods, the correlation was poor. This might partly be because in this species the
baroreflex probably operates also at frequencies below the midfrequency band\textsuperscript{14}. In addition, the pharmacological interventions could have changed the relative influence of non-baroreflex components in the mid-frequency range and consequently changed the phase relation between the blood pressure and the pulse interval in this band. For these reasons, we conclude that the calculation of the transfer gain between the blood pressure and the pulse interval in the mid-frequency band is not suited to measure the BRS in rats. We therefore chose to develop and implement a time-series method. While seeking the best sequential method, we found that the respiratory-related fluctuations seriously hindered the detection of blood pressure ramps lasting longer than five beats in rats. Because the origin of these rapid oscillations is of mechanical rather than of nervous origin in rats\textsuperscript{16,22}, we applied a 10-beat moving-average function to filter out such fast oscillations. In addition this would also facilitate the detection of slower transients in blood pressure and in pulse interval that may be baroreflex-related\textsuperscript{12,14}. Therefore, the focus of our analysis of the spontaneous BRS is restricted to the low-frequency region of the blood pressure and pulse interval fluctuations. Another adaptation of the algorithm was concerned with the shift in beats between the pressure and pulse interval signals. As expected, a one-beat delay between the blood pressure and the heart and pulse interval signals, assuming a latency of the baroreflex of about 260 ms, was inappropriate. To correct this error, we looked for a more appropriate delay of the order of 800 ms\textsuperscript{16,21,2}. However, because the pulse interval is variable, a fixed delay could not be chosen. As a compromise we calculated the BRS at three different delays, three, four and five beats. Averaging these three slopes instead of taking the best fitted
Validation of continuous BRS measurements

slope at delays of three, four or five beats gave the best results. When the delay was not fixed, but rather the BRS was calculated at a variable delay (range 1-12) with the highest correlation coefficient, then, also in SAD and NTS-lesioned rats, a delay could be found at which ramps in blood pressure and pulse interval correlated well. We therefore excluded use of this method because the selection process creates bias towards high BRS estimates in all situations.

It should be noted that the delay of 3-5 beats is probably too short to take account of full sympathetic reactions. Therefore our BRS index is mainly a measure of the parasympathetic reaction. As shown in Figure 3.3 and Table 3.1, the BRS is reduced less by β-blockade than it is by administration of methyl-atropine, indicating the predominance of vagal effects in this measure of the baroreflex sensitivity. We sought to detect possible effects of the sympathetic nervous system by using much longer delays between pressure and pulse interval changes. However, in all cases the results were negative. One of the reasons for this might have been that vagal reactions on the heart may occur superimposed upon slow sympathetically induced changes in heart rate.

In contrast to previously published time-series methods, our algorithm does not select blood-pressure and pulse interval ramps of a certain magnitude. In our method all blood-pressure ramps of four or more beats are included, regardless of the direction and magnitude of the pulse interval change. In cases in which the baroreflex was active, as in intact WKY rats, the average BRS value was always positive. In cases in which the baroreflex had been blocked by pharmacological means or eliminated surgically, a value close to zero was found. These results strengthen our view that baroreflex-mediated variations in pulse interval can be detected on top of all other possible relations between blood pressure and pulse interval changes. These other variations occur in a more random fashion and add only noise to the system. In all circumstances under which the BRS was absent, the detection algorithm returned a gain of zero for the average non-baromodulated triggers. However, in cases in which the baroreflex was active, positive values for the gain were found always. This non-selective method has the advantage that it can be applied also to cases in which it is expected that the baroreflex activity will be absent or very low. When ramps would have been selected on the basis of their magnitude and direction, then only the number of triggers and not so much the slope of the correlation function would fall considerably in cases of low BRS. This introduces the choice of a cut-off point to decide whether the number of triggers is high enough to calculate a value for the BRS. Thus, the present approach differs from previous related methods, in that it does not include a specific gating mechanism that selects only baroreflex blood pressure and pulse interval sequences. Instead of the baroreflex sensitivity in the classical sense, the present method returns estimates about the prevalence of baroreceptor function and indicates the extent to which the baroreceptor functions over time. To track the dynamic baroreceptor function over time it is, however, mandatory to average the individual ramps over time. In our rats, the present algorithm returned about 20 estimates of the baroreflex function each minute. We estimate that a resolution of 1 min is adequate to track the baroreflex function over time. We have used this approach to investigate ultradian variations in BRS in spontaneously hypertensive rats and in WKY rats.

One limitation of our algorithm might be that the absolute BRS values were about
Chapter 3

A factor of three lower than those obtained by the pharmacological method. Several reasons account for this. First, the algorithm calculates the slope as the average of three individual slopes, including slopes with low values. Second, the beat-to-beat fluctuations in pulse interval are relatively higher than those in blood pressure. Therefore, low-pass filtering by the moving average function will affect the pulse interval more and will therefore reduce the slope between these signals, too. Finally, when the algorithm has found a ramp in blood pressure, then all changes in pulse interval are included, regardless of their direction or magnitude. This causes a reduction in the average BRS value, because also nonbaroreflex sequences are included. The reduction in absolute BRS values, however, does not mean that the sensitivity of the method has declined. The BRS index calculated with this on-line method discriminates between WKY rats and spontaneously hypertensive rats and responds to drug treatments. 

It should be borne in mind that the development of the present estimate for the cardiac baroreflex gain from spontaneous fluctuations in blood pressure was done by correlating the performance of the computerized method to the gain derived from pharmacological methods. However, there may be, a priori, limitations to this approach. In the present study the central nervous system was regarded as a system reacting in a linear fashion to changes in input regardless of the magnitude of the input. Although baroreceptor afferents may increase their firing pattern in a linear way within a wide range of pressures, this does not mean that the reaction of the brain to small and large deviations in pressure is similar. With pharmacological methods, the induced changes in blood pressure were much greater than those occurring spontaneously. In the conscious state, the pattern of response to greater input signals could lead to a different or more complete activation-deactivation pattern of brain structures involved in the reflex than when input signals are small. Therefore, the resulting reflex changes in heart rate are not necessarily linear in response to both input levels. This implies that it cannot be decided which approach is superior to the other. We chose to circumvent this problem by matching the pharmacological and spontaneous BRS estimates.

In conclusion, the developed time-series method calculates the gain of the baroreceptor reflex faithfully from spontaneously occurring changes in blood pressure and pulse interval in rats. Furthermore, this method can be implemented on-line in data-acquisition software, making it suitable for long-term continuous measurements of this important physiological control mechanism.

Acknowledgments

We would like to thank Tony Verberne (Department of Pharmacology, Austin Hospital, Heidelberg, Australia) for performing the electrical lesions of the NTS, and Caroline Eerdmans-Tyssen for expert technical assistance.

References

3. Korner P, West MJ, Shaw J, Uther JB: Steady-state properties of the baroreceptor-
Validation of continuous BRS measurements

Chapter 3


Chapter 4
Autonomic control of ultradian and circadian rhythms of blood pressure, heart rate, and baroreflex sensitivity in SHR

Jan Oosting
Harry A.J. Struijker-Boudier
Ben J.A. Janssen

Published in: Journal of Hypertension, 1997, 15:407-410
Chapter 4

Abstract

Objective To examine the influence of the autonomic nervous system on ultradian and circadian rhythms of blood pressure, heart rate and baroreflex sensitivity of heart rate (BRS) in spontaneously hypertensive rats (SHR).

Methods Spontaneous fluctuations in blood pressure, heart rate and BRS in SHR were recorded continuously for 24 h using a computerized system and compared with those in Wistar-Kyoto (WKY) rats. Furthermore, 24 h recordings were performed in SHR during cardiac autonomic blockade by metoprolol and methyl-atropine, vascular autonomic blockade by prazosin, ganglionic blockade by hexamethonium and vagal stimulation by a low dose of scopolamine. The magnitudes of the ultradian fluctuations in blood pressure, heart rate and BRS were assessed by wide-band spectral analysis techniques.

Results The BRS was lower in SHR than in WKY rats throughout the 24 h cycle. In both strains high value were found during the light, resting period, whereas low values were found during the first hours of the dark, active period. The circadian rhythmicity of the blood pressure in SHR was abolished completely during the infusions of prazosin and hexamethonium. In contrast, the circadian rhythmicities of the blood pressure and heart rate were not altered by infusions of metoprolol, methyl-atropine and the low dose of scopolamine. Power spectra of the blood pressure and heart rate lacked predominant peaks at ultradian frequencies and showed 1/f characteristics. In the absence of autonomic tone, the ultradian fluctuations in heart rate, but not in blood pressure, were decreased. The ultradian BRS spectra had no 1/f shape, but showed a major peak at .20 min for 71% of the WKY rats and 42% of the SHR.

Conclusions The influence of the autonomic nervous system on the blood pressure and heart rate in SHR is frequency-dependent. The circadian, but not ultradian, blood pressure rhythmicity is controlled by vascular autonomic activity. Conversely, the circadian, but not ultradian, heart rate rhythmicity is independent of autonomic tone. In rats, just as in humans, the trough in baroreflex sensitivity occurred after the sleeping period, when locomotor activity is resumed.
Introduction

Many studies have examined the role of the sympathetic and parasympathetic nervous systems in the high-frequency oscillations of the blood pressure and heart rate which are related to respiration\textsuperscript{12,13} and cardiovascular reflex mechanisms or Mayer waves\textsuperscript{14-17}. Typically, on these second-to-second time scales, blood pressure and heart rate variabilities show a reciprocal relationship, which is possibly baroreceptor reflex-mediated\textsuperscript{8}. However, less is known about the extent to which the autonomic nervous system contributes to much slower, ultradian fluctuations. Studies in dogs have indicated that the sympathetic nervous system might be involved in blood pressure and heart rate cycles of about 20 min\textsuperscript{9} or of 1-2 h\textsuperscript{10,11}. The exact origin of these cycles is, however, not known. Other studies have found an inverse linear relationship between the logarithm of the frequency and the logarithm of the spectral power of the blood pressure and heart rate\textsuperscript{12,14}. The 1/f characteristic indicates the absence of a single dominant regulatory mechanism and is suggestive of non-linear system characteristics with multiple processes acting on overlapping time scales\textsuperscript{13}.

The 24h diurnal variations in blood pressure and heart rate are thought to be related mainly to behavioural changes, with elevated levels of sympathetic activity during the active period and the dominance of parasympathetic influences during sleep\textsuperscript{16,18}. Now, on this time scale, changes in blood pressure and heart rate occur in parallel. Most evidence for effects of the autonomic nervous system on the 24 h blood pressure and heart rate variability has been derived from studies on essential hypertension\textsuperscript{19}. In humans as well as in experimental animals, differences between daytime and night-time blood pressure and heart rate values are usually more pronounced in hypertensive individuals than they are in normotensive controls\textsuperscript{16,20}. The increased long-term variability generally is explained in terms of elevated sympathetic tone in hypertension or progressively waning control mechanisms\textsuperscript{21}. Treatment with sympatholytic agents can normalize such differences between hypertensive and normotensive states. However, in general, the circadian rhythm in blood pressure and heart rate is preserved\textsuperscript{20,22}. This is not surprising since the doses of those sympatholytic drugs administered usually suppressed the autonomic nervous system only partly. Thus, the extent to which the autonomic nervous system controls the circadian rhythmicity of the blood pressure and heart rate remains unknown.

The present study was designed to evaluate the role played by the autonomic nervous system both in circadian and in ultradian fluctuations in blood pressure and heart rate. Experiments were performed in spontaneously hypertensive rats (SHR) because of the enhanced circadian rhythmicity of their blood pressure and the presumed sympathetic hyperactivity in this model\textsuperscript{17,18,26}. The blood pressure, heart rate and baroreflex sensitivity of the heart rate (BRS) were recorded for 24 h periods during continuous infusions of agents that blocked parts of the autonomic nervous system. In these studies, the BRS was measured continuously by using an algorithm that calculates the gain of the cardiac baroreflex from spontaneous beat-to-beat fluctuations in blood pressure and pulse interval\textsuperscript{23}.
Chapter 4

Methods

Animals

Adult Wistar-Kyōo (WKY) rats and SHR weighing 280-300 g each were used. The rats were obtained from the central Animal Facilities of the Universiteit of Maastricht. Experiments were performed according to the Universiteit Maastricht guidelines. The animals were kept on a 12 h light/12 h dark cycle. After they had undergone surgery, the animals were housed individually in cages and were allowed normal rat chow and drinking water ad libitum.

Surgery

All rats were instrumented with an arterial catheter for measuring their blood pressure and a venous catheter for administration of drugs, as described in detail previously. In short, the catheters were implanted via the femoral vessels and then tunneled through to the lower back of the animal with the animal under pentobarbital anaesthesia. The catheters were exteriorized through a 40 cm long steel spring which was fixed to the back muscles with silicone rubber attached to a piece of mersilene gauze. The venous catheter was filled with a heparinized saline solution (5 U/ml). The steel spring was led to the outside of the cage, where the arterial catheter was connected via a hydraulic swivel (model 375120; Instec Labs, Plymouth Meeting, Pennsylvania, USA) to a low-volume displacement pressure transducer (micro-switch model 156PC 156WL; Honeywell, Inc., Amsterdam, the Netherlands). The arterial canula was kept patent by continuous arterial infusion of heparinized saline solution (30 U/ml) at a rate of 2.4 ml/day. This set-up allowed us to measure blood pressures continuously and to administer drugs without disturbing the rats, for periods lasting usually 4 weeks.

Data acquisition

After the rat had been allowed at least 3 days for recovery the measurements were performed in the home cage of the animal. The arterial pressure transducer was connected to an amplifier that delivered a high-voltage signal to an analogue-to-digital converter board (model 2814; Data Translation, CN Rood, Rijswijk, the Netherlands) connected to an IBM-compatible computer. The blood pressure signal was sampled at 500-1000 Hz. Beat-to-beat pulse interval and mean arterial pressure values were calculated on-line and stored on a hard disc.

Protocols

All experiments were performed in SHR using the following standard protocol. Beat-to-beat recordings were made during intravenous infusions of drugs or saline. The infusions were started 1-4 h before the start of the actual recording. This was done to obtain steady-state pharmacokinetics as well as to avoid acute drug effects within the study period. The infusions were started within the second hour of the dark period. Paired controls were used in these drug studies with at least 2 days between the recording during the saline infusion and that during the drug infusion. The order of the saline infusion and drug infusion was randomized for each animal. The infusion rate in the experiments was 0.2 ml/h.

The following studies were conducted. First, SHR (n = 15) and WKY rats (n = 10)
were compared to study the effect of hypertension. Second, to study the effects of cardiac autonomic blockade, SHR (n = 9) were infused with the β-adrenoceptor antagonist metoprolol (40 mg/kg per 24 h) and the muscarinic antagonist methylatropine (40 mg/kg per 24 h). Third, to block the effects of the autonomic nervous system on the vasculature, SHR (n = 7) were infused with the α-adrenoceptor antagonist prazosin at a dose of 5 mg/kg per 24 h. Fourth, to block the common effenter pathway of the autonomic nervous system, SHR (n = 8) were infused with the ganglion-blocking agent hexamethonium (600 mg/kg per 24 h). Fifth, to stimulate the parasympathetic nervous system, SHR (n = 8) were infused with a low dose of scopolamine. The infusion rate of scopolamine was determined in a pilot experiment with four animals. The infusion rate in this pilot study was increased stepwise from 0.1 to 100 mg/kg per h every 20 min. The optimal dose was considered the dose at which the heart rate decreased the most. This dose (5 mg/kg per h) was used in the 24 h experiments. After the 24 h infusions with hexamethonium and with metoprolol plus methylatropine the efficacy of pharmacological blockade was confirmed by the absence of heart rate changes (< 15 beats/min) in response to injections of nitroglycerin and phenylephrine that changed the blood pressure by -30 to 40 mm Hg. The effectiveness of α-adrenergic blockade by 2.5 mg/kg per 24 h prazosin had been tested in our laboratory previously and found to result in a 160-fold shift towards the right-hand side of phenylephrine-induced increases in blood pressure.24

Data analysis and statistics

The BRS was determined from spontaneous ramps in blood pressure and pulse interval using a modified timeseries method originally used by Bertini et al.25 in cats. The validation and development of an algorithm to assess the cardiac baroreflex gain in rats continuously has been described in detail.23 In essence, the new aspects of this algorithm, in contrast to that used for cats, are that it does not set thresholds for the minimal amplitude of blood pressure or heart rate changes, it does not select the direction of the change in pulse interval and it facilitates the triggering on slower changes occurring in blood pressure over time by low-pass filtering the signals. In short, first, moving averages over 10 beats were calculated both for the blood-pressure signal and for the pulse interval signal to filter out respiration-induced fluctuations. Second, the filtered signal was searched for ramps of decreasing or increasing blood pressure of four beats or more. Third, for each ramp found, the slopes between the blood-pressure ramp and the pulse-interval changes were determined at delays of three, four and five beats. The BRS was calculated as the average value of these three slopes. Average BRS values were stored every minute.

To describe the circadian rhythm in the 24 h experiments, average blood pressure, heart rate and BRS values were calculated every hour. Differences between the averages of the 12 h dark and the 12 h light period were used to characterize the circadian rhythmicity.

The magnitudes of the ultradian fluctuations in blood pressure, heart rate and BRS were analysed by fast Fourier transformation. These parameters were averaged every minute both for the dark period and for the light period. From the 720 data points corresponding to each of these periods, a series of 512 min without missing data was chosen for determination of the power spectrum. The power spectra for the light and
Figure 4.1. Circadian pattern of baroreflex sensitivity (BRS), mean arterial pressure (MAP), and heart rate (HR) in Wistar-Kyoto (O) and spontaneously hypertensive (●) rats. Note the trough in BRS at the transition from the light to dark period. Values are expressed as hourly means ± SEM. The dark period is indicated by the black bars at the base of each plot. Statistical differences are given in Table 4.1.

Figure 4.2. Averaged power spectra of minute-to-minute averages of the baroreflex sensitivity (BRS), mean arterial pressure (MAP), and pulse interval (PI) in Wistar-Kyoto (WKY) and spontaneously hypertensive rats (SHR). Values are expressed as means ± SEM for the upper curve and means - SEM for the lower curve. Note the -9 min and -29 min oscillations of BRS in WKY (△). Average slopes (β ± SD) of the linear regression lines of the MAP and HR spectra are compared between WKY and SHR within each plot. *P < 0.05 versus WKY.

dark periods for each recording were averaged, because in all cases the spectra for the dark, and light periods were very similar. After logarithmic transformation both of the frequency and of the power, linear correlation between the frequency and the power was used to describe the slope and intercept of the power spectrum in the range 0.01–0.5 cycles/min (0.00016–0.0083 Hz).

Values are expressed as means ± SEM unless indicated otherwise. Differences between WKY rats and SHR were assessed using one-way analysis of variance. Effects of pharmacological treatment versus saline control within groups of animals were compared with paired Student’s t-tests. P < 0.05 was considered statistically significant.
Table 4.1. Steady-state values in spontaneously hypertensive rats (SHR) and Wistar-Kyoto (WKY) rats.

<table>
<thead>
<tr>
<th></th>
<th>WKY (n=10)</th>
<th>SHR (n=15)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BRS 24 h (ms/mmHg)</td>
<td>0.36±0.03</td>
<td>0.28±0.02</td>
</tr>
<tr>
<td>BRS Δ Dark-Light (ms/mmHg)</td>
<td>-0.13±0.03</td>
<td>-0.11±0.02</td>
</tr>
<tr>
<td>MAP 24-h (mmHg)</td>
<td>111±1</td>
<td>145±5</td>
</tr>
<tr>
<td>MAP Δ Dark-Light (mmHg)</td>
<td>4.4±0.7</td>
<td>10.9±1.5</td>
</tr>
<tr>
<td>HR 24-h (beats/min)</td>
<td>351±6</td>
<td>309±5</td>
</tr>
<tr>
<td>HR Δ Dark-Light (beats/min)</td>
<td>40±4</td>
<td>44±4</td>
</tr>
</tbody>
</table>

Comparison of 24-h average values of baroreflex sensitivity (BRS), mean arterial pressure (MAP), and heart rate (HR) in SHR and WKY rats. Δ Dark-Light, the difference between average values of the 12 h dark and 12 h light period.: *P < 0.05, versus WKY rats; †P < 0.01, difference between light and dark periods.

Drugs
Metoprolol tartrate (metoprolol), atropine methyl nitrate (methyl-atropine), hexamethonium bromide (hexamethonium), scopolamine hydrochloride (scopolamine) and phentylephrine hydrochloride (phenylephrine) were obtained from Sigma Chemical Company (St Louis, Missouri, USA). Prazosin hydrochloride (prazosin) was obtained from Pfizer (Brussels, Belgium). Sodium nitroprusside (nitroprusside) was obtained from Janssen Chimica (Beerse, Belgium).

Results

SHR versus WKY rats

Figure 4.1 shows the circadian rhythms in blood pressure, heart rate and BRS for SHR and WKY rats. In both strains, the blood pressure and heart rate were higher during the dark (active) period, whereas the BRS was higher during the light (sleeping) period. Average 24 h heart rate and BRS values were lower in SHR than they were in WKY rats, whereas the blood pressure was higher. During the 24h cycle, minimal BRS values were found at the transition from the light to the dark period. This period was accompanied by an increase in blood pressure and heart rate in both strains. The difference in blood pressure between the dark and the light period was smaller for WKY rats than it was for SHR (Table 4.1), whereas such differences in heart rate and BRS were similar.
Table 4.2. Effects of treatment on the 24 h average values and differences between averages for the 12 dark and 12 h light periods (Δ dark-light) of the baroreflex sensitivity (BRS), mean arterial pressure (MAP) and heart rate

<table>
<thead>
<tr>
<th>Group</th>
<th>Methylxanthine</th>
<th>Hexamethonium</th>
<th>Prazosin</th>
<th>Scopolamine</th>
</tr>
</thead>
<tbody>
<tr>
<td>Saline</td>
<td>(n=9)</td>
<td>(n=8)</td>
<td>(n=7)</td>
<td>(n=8)</td>
</tr>
<tr>
<td>BRS 24 h</td>
<td>0.29±0.04</td>
<td>-0.02±0.01 **</td>
<td>0.28±0.02</td>
<td>-0.06±0.02 **</td>
</tr>
<tr>
<td>BRS Δ dark-light</td>
<td>-0.16±0.02 **</td>
<td>0.07±0.01</td>
<td>-0.09±0.03 **</td>
<td>-0.03±0.02</td>
</tr>
<tr>
<td>MAP 24 h</td>
<td>146±5</td>
<td>119±4 **</td>
<td>146±5</td>
<td>137±5</td>
</tr>
<tr>
<td>MAP Δ dark-light</td>
<td>10.3±2.1 **</td>
<td>11.0±3.3 **</td>
<td>13.3±2.9 **</td>
<td>12.9±3.2 *</td>
</tr>
<tr>
<td>HR 24 h</td>
<td>129±5</td>
<td>208±4 **</td>
<td>309±4</td>
<td>308±5</td>
</tr>
<tr>
<td>HR Δ dark-light</td>
<td>44±6 **</td>
<td>36.6±3 **</td>
<td>49±6 **</td>
<td>28±2 **</td>
</tr>
</tbody>
</table>

* P < 0.05, ** P < 0.01, versus saline; † P < 0.05, †† P < 0.01, difference between light and dark periods

The magnitude of the ultradian rhythms was assessed by calculating the power spectra from minute averages. The blood pressure and heart rate spectra lacked predominant peaks. They showed an almost linear relationship in the frequency range 0.01-0.5 cycles per minute (Fig. 4.2). The slope of the linear regression line (β) of the spectrum for blood pressure was greater for SHR than it was for WKY rats [SHR β = 1.43 ± 0.10 (mean ± SD), WKY β = -1.13 ± 0.13; P < 0.01]. Also the slope of the pulse interval spectrum was higher for SHR than it was for WKY rats (SHR β = 1.26 ± 0.08, WKY β = -1.16 ± 0.07; P < 0.05). The magnitude of the ultradian fluctuations in BRS was higher for WKY rats than it was for SHR; however, the difference did not attain statistical significance (P = 0.066). The BRS power spectra did not show a 1/f relation. In the spectra for five of seven WKY rats and five of 12 SHR a small, well-defined peak could be seen with a period in the range 20-25 min. Small peaks were visible at about 9 min in the spectra for two WKY rats and two SHR rats.

Effects of cardiac autonomic blockade

The BRS in SHR during cardiac autonomic blockade with metoprolol and methylxatropine was close to zero throughout the 24 h period (Fig. 4.3). Under this condition the circadian rhythmicity in blood pressure and heart rate was not abolished. Blood pressure and heart rate differences between the dark and the light period were comparable to control differences (Table 4.2). Figure 4.4 shows the power spectra obtained during cardiac blockade with their matched controls. The power spectral density for the blood pressure did not change, but the slope of the power spectrum for the pulse interval became steeper and the value at the intercept decreased (Pulse Interval β<sub>Control</sub> = -1.27 ± 0.08, β<sub>Blockade</sub> = -1.84 ± 0.30; P < 0.05). These changes were due mainly to the
Figure 4.3. Circadian patterns of baroreflex sensitivity (BRS), mean arterial pressure (MAP), and heart rate (HR) in spontaneously hypertensive rats during saline infusion (○, control) and during cardiac autonomic blockade by an infusion of metoprolol and methyl-atropine (●). Values are expressed as hourly means ± SEM. The dark period is indicated by the black bar at the base of each plot. Statistical differences are given in Table I.

Figure 4.4. Power spectra of minute-to-minute averages of mean arterial pressure (MAP), and pulse interval (PI) in spontaneously hypertensive rats during saline infusion (SHR) and during cardiac autonomic blockade by infusion of metoprolol and methyl-atropine (BLK). Average slopes (β±SD) of the linear regression lines of the MAP and PI spectra are compared within each plot. * P<0.05 versus SHR.

Effects of ganglionic blockade

Shortly after the infusion of hexamethonium into SHR had been started, the blood pressure fell considerably (data not shown), but it returned to near-control values within 4 h. Thereafter, the 24 h blood pressure and heart rate averages did not differ from control values, but the baroreflex was blocked (Table 4.2, Fig. 4.5). In the absence of neurogenic tone, the circadian blood pressure rhythmicity was reduced significantly whereas that of the heart rate was maintained. Similarly, as was found during cardiac autonomic blockade, ganglionic blockade did not alter the power spectrum of the blood pressure. However, the spectral density power of the heart rate was reduced greatly.
Figure 4.5. Circadian patterns of baroreflex sensitivity (BRS), mean arterial pressure (MAP), and heart rate (HR) in spontaneously hypertensive rats during saline infusion (○, control) and during ganglionic blockade with hexamethonium (●). Values are expressed as hourly means ± SEM. The dark period is indicated by the black bar at the base of each plot. Statistical differences are given in Table 1.

Figure 4.6. Power spectra of minute-to-minute averages of mean arterial pressure (MAP), and pulse interval (PI) in spontaneously hypertensive rats during saline infusion (SHR) and during infusion of hexamethonium (HEX). Average slopes (β ± SD) of the linear regression lines of the MAP and PI spectra are compared within each plot. * P < 0.05 versus SHR.

This was associated with a slight but significant increase in the slope of the linear regression line of the heart rate (β_{control} = -1.26 ± 0.05, β_{blockade} = -1.52 ± 0.18; P < 0.05) and a decrease in the intercept. These changes were due mainly to a decreased magnitude of the pulse interval fluctuations at the higher frequencies (Fig. 4.6).

Effects of α-adrenergic blockade

Prazosin treatment increased the average 24 h heart rate while the average 24 h blood pressure remained the same (Table 4.2, Fig. 4.7). The 24 h BRS was reduced by 50% (Table 4.2). Prazosin treatment reduced the circadian heart rate and BRS rhythmicities and even reversed the 24 h blood pressure pattern. The power spectrum of the BRS was not altered during prazosin treatment (Fig. 4.8). The spectral density power of the blood pressure was decreased throughout the frequency range and was associated with a slight fall in the linear regression coefficient (β_{control} = 1.44 ± 0.08, β_{prazosin} = -1.23 ± 0.09; P < 0.05). The spectral density power of the heart rate increased and the value at the
Figure 4.7. Circadian patterns of baroreflex sensitivity (BRS), mean arterial pressure (MAP), and heart rate (HR) in spontaneously hypertensive rats during saline infusion (○, control) and during infusion of prazosin (●). Values are expressed as hourly means ± SEM. The dark period is indicated by the black bar at the base of each plot. Statistical differences are given in Table 1.

Figure 4.8. Power spectra of minute-to-minute averages of the baroreflex sensitivity (BRS), mean arterial pressure (MAP), and pulse interval (PI) in spontaneously hypertensive rats during saline infusion (SHR) and during infusion of prazosin (PRA). Average slopes (β±SD) of the linear regression lines of the MAP and PI spectra are compared within each plot. *P<0.05 versus SHR.

The intercept of the linear regression line increased without the slope altering (Fig. 4.8).

Effects of vagal stimulation

During the continuous, low-dose infusion of scopolamine the average 24h heart rate fell by 20 ± 4 beats/min, proving that the vagal stimulation had been maintained throughout this period (Table 4.2). However, under this condition no effects on the blood pressure or BRS were observed. Also the circadian rhythmicities (Fig. 4.9) and ultradian (Fig. 4.10) fluctuations in blood pressure, heart rate and BRS were not altered.
**Figure 4.9.** Circadian pattern of baroreflex sensitivity (BRS), mean arterial pressure (MAP), and heart rate (HR) in SHR during saline infusion (control) and during infusion of low-dose scopolamine. Values are shown as hourly means ± SEM. The dark period is indicated by the black bar at the base of each plot. Statistical differences are given in table 4.1.

**Figure 4.10.** Power spectrum of minute-to-minute averages of baroreflex sensitivity (BRS), mean arterial pressure (MAP), and heart rate (HR) in SHR during saline infusion and during infusion of low-dose scopolamine (SCO). Note the ~9 min oscillation of BRS (Δ). Average slopes (β±SD) of the linear regression lines of the MAP and HR spectra are compared within each plot. * Indicates statistical differences (P<0.05) between control and treated SHR.

**Discussion**

The autonomic nervous system had different, frequency-dependent effects on circadian and ultradian rhythms of the blood pressure, heart rate and BRS. These effects are discussed in detail in separate sections.

**Circadian blood pressure rhythms**

The circadian rhythmicity of the blood pressure was found to depend upon the activity of the autonomic nervous system. The 24 h blood pressure rhythm was suppressed during the infusion of hexamethonium and even reversed by prazosin infusion,
confirming our previous results with this agent. An inversely blood pressure rhythm also occurs in some forms of secondary hypertension and in renin-2 transgenic rats. A possible explanation for these findings is that, under normal physiological conditions, the circadian rhythm of blood pressure is dominated by changes in the sympathetic activity of the vasculature. However, in the absence of neurogenic tone, the circadian rhythms of vasoactive hormones, like vasopressin and renin, which peak during the resting phase of the diurnal cycle, become unmasked. Alternatively, prazosin might have dilated the muscle vascular beds selectively, so that the blood pressure fell more during periods of activity than it did during the resting phase.

**Circadian heart rate rhythms**

In contrast to that of the blood pressure, circadian rhythmicity of the heart rate was found to be largely independent from the autonomic nervous system. During cardiac autonomic blockade with metoprolol and methyl-tropane, the 24 h heart rate pattern was similar to control, although at a lower offset. One may argue that adrenaline mediated $\alpha_{1}$-adrenoceptor stimulation might have determined the 24 h pattern. However, also during ganglionic blockade, the 24 h heart rate rhythm did not differ from the control rhythm. Therefore, our data indicate that intrinsic heart rate is variable throughout the day. The present study, however, does not exclude the possibility that non-nicotinergic nerves play a role. Also in patients, after surgical denervation of the heart after transplantation, circadian rhythmicity of the heart rate is not abolished and parallels physical activity. We therefore suggest that the circadian heart rate rhythm is predominantly caused by physical activity and that atrial filling, which increases as soon as cardiac output rises, plays a major role.

**Circadian rhythms of the BRS**

In this study we found that the 24 h average values of BRS were 23% lower in SHR than they were in WKY rats. Comparable reductions in the baroreflex gain in SHR have been documented before (a reduction by 50% in SHR aged 9-12 weeks; a reduction by 30% in SHR aged 14 weeks). These earlier studies were performed at only one time point of the day. The current study shows that this difference in BRS persists throughout the day. This finding is in accordance with data obtained on humans. Typically, the BRS in SHR showed a circadian rhythm with its highest values occurring during the resting period and minimal values during the first hours of the active period, when blood pressure and heart rate were increasing. An explanation for the trough in BRS during this period could be that the baroreflex has not yet been reset to the increasing pressure and that the rise in heart rate induced by physical activity overrides its action. During the corresponding period in humans there is an increased incidence of cardiovascular incidents such as myocardial infarction. This pattern suggests that the decreased baroreflex gain could be an independent risk factor for these cardiovascular incidents.

The reduction of the BRS in SHR seems to be due mainly to parasympathetic impairment and left ventricular hypertrophy. Therefore we expected that stimulation of the parasympathetic nervous system by infusion of a low dose of scopolamine would increase the BRS. The infusion of scopolamine decreased the average 24 h heart rate by $20 \pm 4$ beats/min. However, the BRS was increased only for a few hours during the dark period. This suggests that scopolamine could have altered the range, but not
Chapter 4

the gain of the baroreflex. Also, the presence of left ventricular hypertrophy per se could have been responsible for such minor effects of scopolamine in SHR. Alternatively, strong vagomimetic effects of scopolamine are possibly only exposed in severely hypersympathetic states, like in heart failure, but not in spontaneous hypertension.

**Ultradian fluctuations in the blood pressure, pulse interval and BRS**

The low-frequency power spectra of the blood pressure and pulse interval showed an almost linear inverse relationship on a double-logarithmic scale within the frequency range 0.01–0.5 cycles/min, without predominant peaks. The slopes of the l/f power spectra both of the blood pressure and of the heart rate were greater for SHR than they were for WKY rats. We have no physiological interpretation of this finding. Wagner et al. found that the slope of the l/f spectrum of the blood pressure in sinu-aortic denervated dogs was higher than that of non-denervated dogs. The increase in the slope could indicate a decreased complexity of the non-linear interactions among the remaining control systems. Alternatively, the increased slope could be due to the enhanced activity of compensatory mechanisms at lower ultradian frequencies. Indeed, after the hyperactivity of the sympathetic nervous system in the sino-aortic denervated dogs had been suppressed with hexamethonium, the slope returned to control values. In the present study in SHR, hexamethonium did not cause such effects. This might indicate that the presumed sympathetic hyperactivity is rather low in adult SHR. In contrast, prazosin did reduce the magnitude of the ultradian fluctuations in blood pressure, without changing its averaged 24 h steady-state level. This suggests that α-adrenergic receptors are involved in the generation of low-frequency fluctuations in blood pressure. Mayer waves in the blood pressure of conscious rats may stem from an intrinsic cyclical activity of resistance arteries with a period of about 10 s. The amplitude of these vasomotor cycles is, amongst other factors, dependent on α-adrenergic stimulation. At present the extent to which such relatively fast fluctuations contribute to the observed ultradian fluctuations in blood pressure is not known. Theoretically, deterministic chaotic control systems with scale invariant or l/f output characteristics can evolve from the interaction of three non-linear equations. Here, we speculate that these non-nervous system-dependent spontaneous vasomotor fluctuations are among the factors contributing to long-term fluctuations in blood pressure.

In contrast to that which was found for blood pressure, the ultradian fluctuations in heart rate are under the predominant control of the autonomic nervous system, since cardiac autonomic blockade with metoprolol plus methyl-atropine and with hexamethonium reduced the spectral power throughout the frequency range. The different effects of the autonomic blocking agents on the blood pressure and heart rate suggest that, at these ultradian frequency ranges, in contrast to higher frequencies, heart rate fluctuations and blood pressure fluctuations are coupled loosely. Whether this feature could result from a reduced baroreceptor feedback in SHR at these frequencies is not known. Wide-band blood pressure fluctuations with periods from 30 s to 30 min in cats were recently shown to be modulated at least partly by the baroreflex. Clearly, more work is needed to unravel the haemodynamic mechanisms involved at these low frequencies.

Ultradian fluctuations in BRS were slightly more pronounced in WKY rats than they were in SHR. Also in WKY rats, rather than in SHR, cycles of ~9 and ~20 min period
24 hour autonomic control of cardiovascular function

in BRS were found. The frequency of the slowest cycle corresponded to that found in blood pressure in dogs after they had been unmasked by cardiopulmonary and sino-
avoicact denervation. The origin of these fluctuations is not known, but we speculate that they are caused by slow oscillations in vascular resistance since such oscillations were absent during prazosin infusion (Fig. 4.8).

One of the limitations of this study is that we cannot exclude the possibility that the administration of the agents to influence the autonomic nervous system might have caused central effects also and thus affected the behavioural activity of the rat. The infusions of hexamethonium and prazosin may have influenced the feeding and drinking behaviour of the rats and the observed cardiovascular effects may thus have been a consequence of these effects, too. Furthermore, the pharmacological blockades by the agents may have varied during the 24 h. Although we recorded the BRS to quantify the functional cardiac blockade with hexamethonium, we cannot exclude the possibility that vascular autonomic blockades was achieved throughout the 24 h period.

Plasma sampling of sympathetic neurotransmitters or vasoactive hormones to monitor the autonomic blockade over time and the endocrine response to it was not possible in the present protocol, since the amount of plasma needed would have disturbed the haemodynamic measurements in the rats. For this reason we cannot estimate the extent to which changes in circulating neurotransmitters and vasoactive hormones may have contributed to the present findings.

In conclusion, the influence of the autonomic nervous system on the blood pressure and heart rate in SHR is frequency-dependent. Circadian but not ultradian blood pressure rhythms are controlled by vascular autonomic activity. The circadian rhythmicity of the heart rate occurs in the absence of autonomic tone, whereas ultradian fluctuations in heart rate depend on it. A trough in BRS occurs for rats, just as it does for humans, after the sleeping period when locomotor activity is resumed.

References

Chapter 4


Chapter 5
Circadian and ultradian control of cardiac output in spontaneous hypertension in rats

Jan Oosting
Harry A.J. Struijker-boudier
Ben J.A. Janssen

Published in American Journal of Physiology 1997; 273 (Heart circ. Physiol. 42): H66-H75
Chapter 5

Abstract

The aim of the study was to test whether circadian and ultradian variations of cardiac output (CO) in spontaneously hypertensive rats (SHR) differ from those in normotensive Wistar Kyoto rats (WKY). Twenty-four-hour beat-to-beat recordings of CO (by electromagnetic flow probe) and mean arterial pressure (MAP) were performed in the absence and presence of cardiac autonomic blockade with metoprolol and atropine methylnitrate. Ultradian variability was analyzed by spectral analysis on beat-to-beat data series (high-frequency range) and on averaged minute-to-minute data series (low-frequency range). In general, circadian and ultradian rhythms of CO were similar in SHR (n = 10) and WKY (n = 9). Values of CO were high during the dark and low during the light period, whereas total peripheral resistance was highest during the light period. During cardiac autonomic blockade, relative differences between averaged values of CO over the dark and light periods were reduced. High-frequency spectral power of CO was mainly confined to fluctuations related to respiration and was not influenced by cardiac autonomic blockade. At low-frequency ranges, power spectra of CO lacked a dominant oscillator but showed 1/f characteristics. During cardiac autonomic blockade, low-frequency spectral power of CO fell without changing the 1/f characteristics. These findings suggest that dynamic control of CO is not altered in SHR and that autonomic effects on CO are frequency dependent. In most frequency ranges, the relative variation of CO was higher than that of MAP. Thus, over 24 h in both adult SHR and WKY, MAP is controlled within a more narrow range than CO.
Introduction

Arterial blood pressure fluctuates diurnally with low values during periods of sleep and high values during periods of activity. The circadian patterns of cardiac output (CO) and peripheral resistance, which are responsible for the diurnal variation in arterial pressure, are less well known. Only in a few studies in humans, nonhuman primates, dogs, and rats, has CO, or indirect measures of it, been recorded over a full diurnal cycle. In these studies, CO was low and peripheral resistance was high during periods of sleep, whereas the reverse pattern was observed during periods of activity. The evidence thus far suggests that the diurnal profile of CO is closely linked to the physical activity pattern. The diurnal changes in peripheral resistance, especially during periods of sleep, when metabolic demand and sympathetic tone are low, are probably governed by autoregulatory processes. Except for one study in cardiac transplant recipients, all aforementioned studies were confined to healthy volunteers and healthy animals.

In the established phase of spontaneous hypertension, both in humans and animal models, CO is normal, whereas peripheral resistance is elevated. These results were obtained mainly from measurements performed over relatively short periods of time and mostly in resting conditions. However, circadian fluctuations in CO may be elevated in spontaneous hypertension and contribute to the increased diurnal fluctuations of blood pressure associated with this disease in humans as well as in animal models. In addition, ultradian (<24 h) fluctuations in CO output could also be different in spontaneous hypertension. Indicative for this is the finding that short-term variability of heart rate (HR) is decreased. Generally, this is explained as being caused by impaired vagal control of HR. Whether this holds true for the regulation of CO in hypertension is not known.

The first aim of the present study was to investigate the circadian and ultradian profiles of systemic hemodynamic parameters in spontaneous hypertension. The second aim was to examine the dynamic interaction of CO, arterial pressure, and peripheral resistance during a full circadian cycle. To this end, we recorded CO and mean arterial pressure (MAP) on a beat-to-beat basis for 24 h in spontaneously hypertensive rats (SHR) and Wistar-Kyoto rats (WKY). Experiments were performed under normal conditions as well as during cardiac autonomic blockade with metoprolol and atropine methyl nitrate. The dynamic relationship between blood pressure, CO, and peripheral resistance was examined by spectral analysis.

Methods

Animals

Adult male WKY and SHR rats weighing 280-300 g were used. The rats were obtained from the Central Animal Facilities of the Universiteit Maastricht. Experiments were performed according to institutional guidelines and approved by the institutional ethics committee. The animals were kept on a 12:12-h light-dark cycle. After surgery, the animals were housed individually in cages and were allowed normal rat chow and drinking water ad libitum.
Chapter 5

Surgery

All rats were instrumented with an electromagnetic (EM) flow probe around the ascending aorta, an arterial catheter for measurement of blood pressure, and a venous catheter for administration of drugs. Surgery was performed in two stages, both under pentobarbital sodium (60 mg/kg ip) anesthesia and sterile conditions. In the first stage, the EM flow probe was placed. The trachea was intubated (PE-240). The skin and muscles overlying the third right intercostal spaces were separated and cut. After respiration was started with room air (60 strokes/min, tidal volume 3 ml), the thorax was opened. The ribs were spread with a small retractor, and the aorta was dissected free. The EM flow probe (2.7 mm; Skalar, Delft, The Netherlands) was placed around the ascending aorta just above the heart. The thorax was closed by pulling the ribs together with 3-0 silk, and the cable of the probe was fixed to the ribs. Muscles were sutured separately with 5-0 silk. The cable was guided to the neck, where the connector was fixed to the muscles and skin. The skin was sutured (3-0 silk), and subatmospheric intrathoracic pressure was restored by insertion of a Silastic drain through a stab wound between the sixth and seventh ribs and application of a negative pressure of 10 cmH2O. The animals were allowed 2-3 days of recovery before the second stage of surgery. In this latter procedure, the abdominal aorta and the inferior vena cava were cannulated via the femoral vessels. The catheters were tunneled subcutaneously to the neck, where they were exteriorized. The catheters were filled with a heparinized saline solution (5 U/ml).

Protocol and data acquisition

The measurements were performed with the rats being kept in their home cage. Two to three days after implantation of the catheters, the rats were connected to the equipment. The cable of the flow probe and the arterial and venous catheters were guided through a rubber tube to protect them. The rubber tube was attached to a balanced, freely moving arm above the rat to allow the animal full movement. The EM flow probe was connected to a sine-wave EM flowmeter (Transflow 601 system, type MDL1401, Skalar). The arterial catheter was connected to a low-volume displacement pressure transducer (microswitch, model 156PC 156WL, Honeywell, Amsterdam, The Netherlands) that was connected to an amplifier that delivered a high-voltage signal to an analog-to-digital converter board (model 2814, Data Translation, CN Rood, Rijswijk, The Netherlands) mounted in an IBM-compatible computer. Both the flow and pressure signals were sampled at 1,000 Hz. A dedicated computer program was used to identify on-line the end-diastolic, flow of each beat, which was taken to be zero. Then, from beat to beat, the pulse interval (PI), HR, MAP, stroke volume (SV), CO, and total peripheral resistance (TPR) were calculated and stored on hard disk. At least 24 h were allowed to pass before measurements began, to accustom the animal to the experimental conditions.

Measurements were made for two consecutive days. An intravenous infusion of a 0.9% saline solution (0.2 ml/h) was given on the first day as vehicle. The total volume infused (5 ml/24 h) is low compared with average water intake (~30 ml/24-h) in rats. On the second recording day, the cardiac autonomic system was blocked by an infusion of metoprolol (1.65 mg.kg\(^{-1}\).h\(^{-1}\)) and atropine methyl nitrate (1.65 mg.kg\(^{-1}\).h\(^{-1}\)) dissolved in saline at a rate of 0.2 ml/h. To obtain steady-state pharmacokinetics and to avoid acute
drug effects, the recording was started 2 h after the infusion had started. The saline
infusions were started in the second hour of the dark period. The recording during
cardiac autonomic blockade was started in the fourth or fifth hour of the dark period. At
the end of the 24-h infusion period, cardiac autonomic blockade was verified by recording
HR changes after intravenous bolus injections of sodium nitroprusside and phi-
nyephrine. After 24-h. infusion of metoprolol and atropine methylxantrate, the
phenylephrine-induced changes were MAP +40 ± 6 mmHg and HR -6 ± 1 min⁻¹ in SHR
(n = 10) and MAP +40 ± 2 mmHg and HR -5 ± 1 min⁻¹ in WKY (n = 9). Nitro-
prusside-induced changes were MAP -26 ± 2 mmHg and HR +9±2 min⁻¹ in SHR and
MAP -26±2 mmHg and HR +11±2 min⁻¹ in WKY. In intact WKY and SHR, such pressure
changes evoke HR changes >50 min⁻¹, confirming a functional autonomic cardiac
blockade. The fixed order of recording first vehicle and then treatment was chosen for
a technical reason. In this particular setup, we found that when rats were tethered for
longer periods of time the chance of dysfunction of the flow probe increased con-
siderably. Therefore, when the reversed order would have been done, allowing for full recov-
er of the drug effects, the study would have been prolonged by 3 days.

Data analysis

Data were included for analysis only when both the 24-h recordings during saline
and cardiac autonomic blockade were completed. To construct the circadian patterns,
the beat-to-beat values of CO, MAP, HR, peripheral resistance, and SV were averaged
over every hour of the recording. Coefficients of variation were calculated as 100 times
the standard deviation of all 24-h values divided by the 24-h average value. As a
measure of circadian rhythmicity, the relative circadian amplitude of each parameter
was calculated as the difference between the maximum and minimum values of the hourly
averages in a 24-h period normalized by the average 24-h value. In addition, we calcu-
lated the relative difference between the average values over the 12-h dark period and
the 12-h. light period normalized to the average 24-h values.

Spectral analysis in high-frequency range. The dynamic relationship between
the systemic hemodynamic parameters was analyzed by spectral analyses as described
previously16. For the analysis in a high-frequency (HF) range, a stable period of 6,000
beats (15-20 min) was selected from the third hour of the dark period and the third hour
of the light period. For spectral analysis, equidistantly sampled data are required.
Therefore, from these two series of beat-to-beat data, equidistant data series were constructed
using an algorithm that extracted from sequential 200-ms time windows the maximum
value of each parameter. Because PI was never >200 ms, the width of the time window
was set at 200 ms to avoid aliasing. Spectral density was calculated by a fast Fourier
transform on consecutive blocks of 1,024 values, and the results were averaged for all
blocks in a recording. In this way we were able to explore the frequency range of 0.05-2.5
Hz with a resolution of 0.005 Hz. Data were averaged over a low-frequency (LF; 0.05-0.25
Hz), a midfrequency (MF; 0.25-0.55 Hz), and an HF (1.0-2.1 Hz) band. In these selected
frequency ranges, mechanisms related to autoregulation (LF), the autonomic nervous
system (MF), and the respiratory cycle (HF) have been identified16.

Spectral analysis in LF range. For the analysis in the LF range, values of CO,
blood pressure, and peripheral resistance were averaged every minute for both the dark
and light periods. From the 720 data points in each of these periods, a series of 512 min
was taken for the calculation of the power spectrum. Because for all rats power spectral density in the light and dark periods was similar, data were averaged over these periods. After logarithmic transformation of both the frequency (f) and the power, a linear correlation between f and power was used to describe the slope (β) and intercept of the linear part of the power spectrum between 0.01 and 0.5 cycles/min. The inverse relation between f and spectral density power reflects the operation of multiple processes with overlapping time scales.\(^{17}\) The slope of the 1/f spectrum is independent of total power and is an index of the complexity of the system controlling the parameter. This index has recently been useful in research describing changes in the labile behavior of HR and blood pressure.\(^{18,31}\)

**Transfer function analysis.** In addition to spectral density power, the transfer function was calculated, with CO as the input signal and arterial pressure as the output signal.\(^{14}\) The transfer gain was normalized by the average steady-state resistance. In this case, the normalized gain is the ratio of the fractional variation in pressure to CO. A value of 1 for the gain indicates that, when the input (CO) fluctuates by some fraction of the mean value, the output (arterial pressure) undergoes the same fractional variation, as would be the case in a rigid vascular bed, without resistance variation. A value <1 signifies that the fractional variations in pressure are smaller than in CO. A gain >1 indicates that relative fluctuations in pressure are greater than fluctuations in CO. In physiological terms, the latter case means that the system prefers to control flow rather than pressure, which would be indicative of autoregulatory behavior. In addition, we computed the phase and coherence. The phase indicates the temporal relationship between the CO and blood pressure signals in the frequency domain. The coherence can have a value between 0 and 1 and is a frequency-domain estimate of the degree that the blood pressure variance can be explained by a linear operation of the CO variance.

**Two-dimensional frequency distributions.** Mutual interactions between systemic hemodynamic parameters were examined by contour plots. These plots, comparable to topographical altitude maps, were made by calculating two-dimensional frequency histograms and plotting lines through bins of 30° with equal numbers of occurrences. The scales of the axes were normalized to the average 24-h value. The advantage of such plots over time-course graphics is that they 1) illustrate in more detail the range of the spontaneous variations of parameters and 2) reveal clusters of preferred combinations between parameters that are not visible in time-course graphics. We have used these techniques to illustrate our modified view on the concept of a set point and the concept of autoregulation.\(^{21,22}\) Preferred combinations of parameters were identified and counted as such when the altitude lines enclosed a combination that occurred at least 600 times or more than surrounding combinations.

**Statistics**

Data are presented as means ± SE. To compare effects of cardiac autonomic blockade versus saline in SHR and WKY, a two-way analysis of variance was used with SHR versus WKY as a between factor and saline versus autonomic blockade as a within factor. Statistical significance was accepted at P < 0.05.
Table 5.1. Average 24-h values and lability indices of systemic hemodynamics

<table>
<thead>
<tr>
<th></th>
<th>WKY (n=9)</th>
<th></th>
<th>SHR (n=10)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control</td>
<td>Blockade</td>
<td>Control</td>
<td>Blockade</td>
</tr>
<tr>
<td><strong>1A: Average 24-h values</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MAP (mmHg)</td>
<td>111±3</td>
<td>99±3 $</td>
<td>135±4 *</td>
<td>121±4 $*</td>
</tr>
<tr>
<td>CO (ml/min)</td>
<td>77±4</td>
<td>71±3 $</td>
<td>76±2</td>
<td>66±2 $</td>
</tr>
<tr>
<td>TPR (mmHg/ml/min)</td>
<td>1.51±0.09</td>
<td>1.45±0.10</td>
<td>1.83±0.10 *</td>
<td>1.89±0.10 *</td>
</tr>
<tr>
<td>HR (beats/min)</td>
<td>351±6</td>
<td>333±7 $</td>
<td>345±4</td>
<td>317±4 $*</td>
</tr>
<tr>
<td>SV (μl)</td>
<td>220±11</td>
<td>213±11</td>
<td>221±6</td>
<td>207±7 $</td>
</tr>
<tr>
<td><strong>1B: Coefficients of variation (of all 24-h beats)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MAP (%)</td>
<td>8.7±0.6</td>
<td>13.0±2.9 $</td>
<td>8.9±0.7</td>
<td>12.1±1.1 $</td>
</tr>
<tr>
<td>CO (%)</td>
<td>14.1±0.7 †</td>
<td>12.1±1.0 †</td>
<td>12.5±0.5  †</td>
<td>11.7±0.9  †</td>
</tr>
<tr>
<td>TPR (%)</td>
<td>17.1±1.2 †</td>
<td>19.5±2.4</td>
<td>15.1±1.1  †</td>
<td>17.4±1.4  †</td>
</tr>
<tr>
<td>HR (%)</td>
<td>11.4±0.5</td>
<td>5.6±0.3 $</td>
<td>11.5±0.5</td>
<td>5.6±0.7 $</td>
</tr>
<tr>
<td>SV (%)</td>
<td>13.0±0.6</td>
<td>11.7±0.8</td>
<td>11.5±0.4 *</td>
<td>10.7±0.7</td>
</tr>
<tr>
<td><strong>1C: Relative circadian amplitudes</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MAP (%)</td>
<td>16.8±2.2</td>
<td>34.1±8.7</td>
<td>20.3±2.2</td>
<td>26.8±2.9</td>
</tr>
<tr>
<td>CO (%)</td>
<td>30.1±3.2 †</td>
<td>28.7±4.4</td>
<td>28.0±1.8  †</td>
<td>26.4±3.7</td>
</tr>
<tr>
<td>TPR (%)</td>
<td>33.7±3.8 †</td>
<td>45.4±8.2</td>
<td>33.5±4.0 †</td>
<td>38.9±6.9</td>
</tr>
<tr>
<td>HR (%)</td>
<td>23.3±1.9</td>
<td>16.8±1.4 $</td>
<td>26.9±2.7</td>
<td>16.1±2.0 $</td>
</tr>
<tr>
<td>SV (%)</td>
<td>24.8±2.7</td>
<td>21.4±2.1</td>
<td>22.4±2.0</td>
<td>18.7±2.7</td>
</tr>
<tr>
<td><strong>1D: Relative difference between 12-h dark period and 12-h light period</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MAP (%)</td>
<td>0.1±1.7</td>
<td>7.6±4.8</td>
<td>0.4±1.5</td>
<td>4.0±2.5</td>
</tr>
<tr>
<td>CO (%)</td>
<td>11.3±2.2 †</td>
<td>7.8±2.5 $</td>
<td>10.6±1.2 †</td>
<td>5.5±2.3 $</td>
</tr>
<tr>
<td>TPR (%)</td>
<td>-11.5±2.5 †</td>
<td>0.2±5.7</td>
<td>-10.0±1.5 †</td>
<td>1.7±4.1 $</td>
</tr>
<tr>
<td>HR (%)</td>
<td>4.7±1.9</td>
<td>6.1±1.3</td>
<td>9.9±1.9</td>
<td>4.7±1.6 $</td>
</tr>
<tr>
<td>SV (%)</td>
<td>6.6±1.9</td>
<td>1.6±2.3</td>
<td>0.7±1.3 *</td>
<td>0.8±1.5 $</td>
</tr>
</tbody>
</table>

Mean values and 24-h lability indices of mean arterial pressure (MAP), cardiac output (CO), total peripheral resistance (TPR), heart rate (HR), and stroke volume (SV) in WKY and SHR during normal conditions (control) and during cardiac autonomic blockade with metoprolol + methyl-ATropine (blockade). Statistical differences between and within groups (ANOVA) are indicated as: * p (WKY, SHR) < 0.05, $ p (control, blockade) < 0.05, and † p (MAP, CO) or p (MAP, TPR) < 0.05.
Chapter 5

Figure 5.1. Average hourly hemodynamic pattern of 24-h recordings in WKY (n=9) and SHR (n=10) during saline infusion. The saline infusion were started between 13-14 h. Cardiac output (CO), mean arterial pressure (MAP), heart rate (HR), total peripheral resistance (TPR), and stroke volume (SV) are shown. The dark period is indicated by the black bar on the time axis. Statistical differences between SHR and WKY in terms of average values, coefficient of variation and relative circadian amplitude are given in Table 1.

Figure 5.2. Average hourly hemodynamic pattern of 24-h recordings in the same animals as in Figure 1 during cardiac autonomic blockade. The recordings during the infusions of metoprolol + methyl-arozone were started between 16-18 h. Cardiac output (CO), mean arterial pressure (MAP), heart rate (HR), total peripheral resistance (TPR), and stroke volume (SV) are shown. The dark period is indicated by the black bar on the time axis. Statistical differences between SHR and WKY as well as between control and blockade are given in Table 1.

Results

Steady state systemic hemodynamics

In SHR, average 24-h MAP and peripheral resistance were higher than in WKY (Fig. 5.1, Table 5.1). No difference was found in average 24-h HR, CO, and SV In both SHR and WKY, CO and HR exhibited higher values during the dark, active period, whereas peripheral resistance showed higher values during the light, resting period. This was quantified by calculating the relative difference between the average values over the dark and light periods (see Table 5.1).

The 24-h coefficients of variation of the systemic hemodynamic parameters were similar for SHR and WKY (see Table 5.1). In both strains, the coefficient of variation of
Circadian control of cardiac output

Table 5.2: Relative spectral density power of systemic hemodynamics in 3 frequency ranges

<table>
<thead>
<tr>
<th>Frequency range</th>
<th>WKY (n=9)</th>
<th>SHR (n=10)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control</td>
<td>Blockade</td>
</tr>
<tr>
<td>MAP (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LF</td>
<td>35.9±2.0</td>
<td>34.5±8.9</td>
</tr>
<tr>
<td>MF</td>
<td>7.2±1.0</td>
<td>5.9±1.0</td>
</tr>
<tr>
<td>HF</td>
<td>1.5±0.2</td>
<td>2.5±0.7</td>
</tr>
<tr>
<td>CO (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LF</td>
<td>9.8±0.6</td>
<td>8.9±0.3</td>
</tr>
<tr>
<td>MF</td>
<td>8.6±0.5</td>
<td>8.1±0.3</td>
</tr>
<tr>
<td>HF</td>
<td>40.9±1.4</td>
<td>47.2±1.5*</td>
</tr>
<tr>
<td>TPR (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LF</td>
<td>11.5±0.6</td>
<td>13.4±0.8</td>
</tr>
<tr>
<td>MF</td>
<td>8.6±0.7</td>
<td>8.4±0.4</td>
</tr>
<tr>
<td>HF</td>
<td>34.8±1.6</td>
<td>39.4±1.7</td>
</tr>
</tbody>
</table>

Relative spectral density power of mean arterial pressure (MAP), cardiac output (CO), and total peripheral resistance (TPR), in 3 frequency ranges: Low frequency (LF) = 0.05 - 0.25 Hz; Mid Frequency (MF) = 0.25 - 0.55 Hz; High Frequency (HF) = 1.0 - 2.1 Hz. Statistical differences between and within groups (ANOVA) are indicated as: * p (WKY, SHR) < 0.05, $ p (control, blockade) < 0.05

Blood pressure was significantly smaller than those of CO and peripheral resistance. The relative circadian amplitudes of the systemic hemodynamic parameters (summarized in Table 5.1) were, in general, 20-30% of the average 24-h mean value. These values did not differ between SHR and WKY. However, in both groups, the relative circadian amplitude of blood pressure was significantly smaller than the relative amplitudes of CO and peripheral resistance.

Cardiac autonomic blockade produced comparable systemic hemodynamic effects in WKY and SHR (Fig. 5.2, Table 5.1). Average 24-h values for CO, MAP, and HR decreased significantly, whereas peripheral resistance did not change. In SHR, the average 24-h value for SV fell slightly but significantly, whereas SV remained unchanged in WKY (Table 5.1). On autonomic blockade, the coefficient of variation for HR decreased, whereas that for blood pressure increased (Table 5.1). The coefficients of variation for CO, peripheral resistance, and SV remained unaltered. The effects of cardiac autonomic blockade on the 24-h rhythmicity of the systemic hemodynamic parameters are indicated in Table 5.1. In both SHR and WKY, the relative circadian amplitude of HR fell during cardiac autonomic blockade, whereas the 24-h rhythmicity of all other parameters was unchanged. In terms of relative dark-light differences, 24-h rhythmicity of CO fell significantly in both WKY and SHR during cardiac autonomic blockade. In addition, in SHR but not in WKY, the relative differences between the dark and light period averages of HR and TPR were reduced.

Figures 5.1 and 5.2 show that during control as well as during cardiac autonomic blockade CO, arterial pressure, and HR rose in the beginning of the dark, active period. At the start of the light period CO decreased, which was due to a fall in HR. SV declined
Chapter 5

Figure 5.3. Averaged power spectra of systemic hemodynamics at high frequency ranges. Mean arterial pressure (MAP), cardiac output (CO), and total peripheral resistance (TPR) are shown. The left panels show WKY (n=9, dotted lines) and SHR (n=10, continuous lines) during saline infusion. The right panels show the results obtained in the same animals during cardiac autonomic blockade. Statistical differences in selected frequency bands are presented in Table 2. The range of the low (LF), mid (MF), and high (HF) frequency bands is indicated by bars at the top of the figure: (LF: 0.05-0.25 Hz, MF: 0.25-0.55 Hz, and HF: 1.0-2.1 Hz).

Figure 5.4. Averaged transfer gain, phase, and coherence between cardiac output and blood pressure at high frequency ranges. The left panels show WKY (n=9, dotted lines) and SHR (n=10, continuous lines) during saline infusion. The right panels show the results obtained in the same animals during cardiac autonomic blockade. The range of the low (LF), mid (MF), and high (HF) frequency bands is indicated by bars at the top of the figure: (LF: 0.05-0.25 Hz, MF: 0.25-0.55 Hz, and HF: 1.0-2.1 Hz).

progressively during the light period. Peripheral resistance was highest during the sleeping period and fell during the transition from the light to the dark period.

HF dynamics

Average HF spectral density power of CO, arterial pressure, and peripheral resistance is presented in Fig. 5.3. In both SHR and WKY, fluctuations of CO were mainly found at the respiratory-related peaks, which were located at slightly lower frequencies in SHR than in WKY. In contrast to the spectra of CO and peripheral resistance, spectral density power of arterial pressure lacked clear respiratory-related peaks and was mostly confined to frequencies <0.5 Hz. The relative powers of CO, arterial pressure, and peripheral resistance are compared between SHR and WKY in three selected frequency bands in Table 5.2. During control conditions as well as cardiac autonomic blockade, the MF power of arterial pressure, but not of CO and peripheral resistance, was significantly greater (factor of ~2) in SHR than in WKY. The phase relationship between CO and arterial pressure showed a transition from positive to negative values at 0.4 Hz.
Figure 5.5. Averaged double logarithmic power spectra of systemic hemodynamics at low frequency ranges. Mean arterial pressure (MAP), cardiac output (CO), and total peripheral resistance (TPR) are shown. The left panels show WKY (n=9, dotted lines) and SHR (n=10, continuous lines) during saline infusion. The right panels show the results obtained in the same animals during cardiac autonomic blockade. The slopes of the linear regression lines of the power spectra, which were calculated between 0.5 and 0.01 cycles/min, are indicated as $\beta$. Values are means SD. * indicates a significant difference (p<0.05) between control and blockade. Differences between SHR and WKY were not statistically significant.

Figure 5.6. Averaged transfer gain, phase, and coherence between cardiac output and blood pressure at low frequency ranges. The left panels show WKY (n=9, dotted lines) and SHR (n=10, continuous lines) during saline infusion. The right panels show the results obtained in the same animals during cardiac autonomic blockade. Differences between SHR and WKY were not statistically significant.

indicating that MF fluctuations in pressure were leading those in CO beyond 0.4 Hz (Fig. 5.4). The normalized transfer gain between CO and arterial pressure was < 1 over the frequency range from 0.1 to 2.5 Hz (Fig. 5.4). Thus, in this frequency range, fluctuations of CO were relatively greater than those of blood pressure.

**LF dynamics**

The spectra of blood pressure, CO, and peripheral resistance in the LF range lacked predominant peaks, and total power was not different between SHR and WKY. When both frequency and power were logarithmically transformed, CO and arterial pressure showed an almost linear relationship with frequency in the range from 0.01 to 0.5 cycles/min (Fig. 5.5). The slopes of these linear regression lines are indicated in Fig. 5.5 and were not different between SHR and WKY. Cardiac autonomic blockade reduced spectral density power of CO but not of arterial pressure and peripheral resistance. Under this condition, the slope of the regression lines of peripheral resistance spectra increased in both SHR and WKY. Under control conditions, the normalized transfer
Figure 5.7. Contour plots showing the 24-h range of interaction of beat-to-beat recorded systemic hemodynamics during intravenous infusion of saline and metoprolol + methyldigoxin (blockade) in a representative WKY and SHR. For each animal the contour plots (birdseye view of 2-dimensional frequency distribution) between mean arterial pressure (MAP) and pulse interval (PI), MAP and cardiac output (CO), MAP and total peripheral resistance (TPR), and CO and TPR are given. Each plot contains about 300,000 data points. All parameters are plotted relatively to their 24-h average value, with the first-mentioned parameter on the abscissa and the second on the ordinate. Altitude lines are plotted with intervals of 300. In the MAP & PI plot combinations mostly showed 2 or more preferred states. The MAP & CO, MAP & TPR, and CO & TPR plots show only a single preferred state. Note the relative difference in variation of CO and MAP and the inverse relationship between CO & TPR.
gain between CO and blood pressure was close to 1 in both SHR and WKY (Fig. 5.6), and the phase was close to 0. Thus, in this frequency range, the magnitude of the variations in CO were comparable to those in blood pressure and occurred concomitantly. During cardiac autonomic blockade the transfer gain between CO and blood pressure increased to ~1.5 in both rat strains, which was due to the decreased fluctuations in CO. The phase relation did not change.

**Two-dimensional frequency distributions**

Contour plots of the two-dimensional frequency distributions of the systemic hemodynamics are presented for a representative WKY and a representative SHR in Fig. 5.7. Under control conditions, the contour plots of MAP and PI showed multiple peaks in 8 of 10 SHR and 7 of 9 WKY. The average number of peaks in these plots was 2.5 ± 0.3 in SHR and 1.9 ± 0.2 in WKY. In two SHR and one WKY, the peaks seemed to be located around a center of relatively few occurrences.

The contour plots of MAP and CO, MAP and TPR, and CO and TPR displayed, in general, a single peak. In 3 of 10 SHR and 2 of 9 WKY a second peak was found. The average number of peaks in these three plots was 1.4 ± 0.2 in SHR and 1.2 ± 0.1 in WKY. The blood pressure value found at the most frequent combination in such plots generally coincided with the blood pressure value at the highest peak in the MAP and PI plots. The elongated shape of the MAP and CO plots indicates that over the 24-h period, the relative variability of CO was higher than the relative variability of blood pressure. A similar pattern occurred when arterial pressure was plotted versus peripheral resistance. In contrast, the frequency distributions of CO and TPR appeared to be inversely related.

After cardiac autonomic blockade, multiple peaks appeared in all contour plots between MAP and PI. The average number of preferred combinations increased significantly to 3.6 ± 0.3 in SHR and 3.1 ± 0.3 in WKY. It was apparent that the overall relative variability of PI was decreased (see also Table 5.1). The most frequent value of blood pressure was often lower than its average 24-h mean value. During cardiac autonomic blockade, the contour plots relating MAP and CO, MAP and TPR, and CO and TPR exhibited multiple peaks in 3 of 10 WKY and 2 of 10 SHR. The average value was not different from control and was 1.1 ± 0.1 in SHR and 1.3 ± 0.2 in WKY. In comparison with the control plots, the figures show that the range of the variation in CO was hardly changed, whereas the range in arterial pressure was greater. This is in agreement with Table 5.1, which shows that the 24-h coefficient of variation of blood pressure is increased during blockade.

**Discussion**

The main new findings of this study are: 1) circadian rhythmicity of CO is similar in adult SHR and WKY; 2) the magnitude of the circadian rhythmicity of CO is reduced by blockade of the cardiac autonomic nervous system; 3) short-term dynamic behavior of CO consists mainly of mechanically induced fluctuations related to respiration; 4) at LF ranges, power spectra of CO lack a dominant oscillator but show 1/f characteristics; and 5) over 24-h periods in both SHR and WKY, variability of CO is relatively greater.
than variability of blood pressure. In these time frames, control of blood pressure is preferred over control of CO.

**Circadian rhythmicity of systemic hemodynamics**

In agreement with previous studies, at least in absolute values, the circadian rhythm of MAP was more pronounced in SHR than in WKY. However, coefficients of variation and relative circadian amplitudes of blood pressure were not different between the strains. This suggests that the systems involved in the regulation of the circadian blood pressure rhythm are not different between strains but that the difference is merely a consequence of the elevated pressure. It was shown previously in SHR that the elevated pressure is not a result of increased CO. We now also demonstrate that the circadian fluctuations of CO, in both absolute and relative terms, are similar in adult WKY and SHR. CO was highest during the dark, active period of these nocturnal animals and lowest during the light, sleeping period. Thus, over the whole 24-h cycle, increased peripheral resistance is responsible for the high blood pressure. In SHR and WKY, the diurnal patterns of CO and peripheral resistance are inversely related. Peripheral resistance was low during the dark, active period and high during the light, sleeping period.

The hemodynamic mechanisms underlying the inverse relationship of the diurnal patterns of CO and peripheral resistance are not fully clear. In the present study, the fall of CO during the light phase was mediated by reductions of HR as well as SV in both SHR and WKY. During cardiac autonomic blockade, values of CO were also higher during the dark period than during the light period, although the relative differences between these periods were significantly attenuated. This suggests that both nervous and nonnervous factors are involved in the generation of the 24-h rhythm of CO. However, as indicated by Table 5.1 and the contour plots in Fig. 5.7, the range of the variation of CO with circadian period is not diminished by cardiac autonomic blockade. Cardiac autonomic blockade reduced specifically the amplitude of the 24-h cycle of CO but not cycles of shorter periods. As depicted in Figs. 5.1 and 5.2, at the beginning of the light period, CO fell because of an abrupt fall in HR. SV declined progressively in this period and was lowest at the end of the light period, when HR was rising again. Similar observations were obtained by Smith et al. in normotensive Wistar rats. In humans and nonhuman primates, a reduction in HR seems the dominant factor involved in the fall of CO at sleep, with only minor changes in SV. Such species differences could be due to differences in posture. Rats remain in the same position, whereas humans and nonhuman primates assume a recumbent position at sleep. However, there is redundancy in this control system. When HR was prevented from falling overnight by atrial demand pacing, CO fell in nonhuman primates by a reduction in SV. The reduction in CO during sleep may be due to venous pooling of the blood or to a reduction in plasma volume. The slow, progressive fall of SV during the light period in rats could be indicative of the latter mechanism. Rats do not drink during the light period, and water volume may decline with ongoing urine formation, expiration of air, or perspiration. Interestingly, these diurnal changes in hemodynamics and plasma volume could be governed by increased levels of atrial natriuretic peptides, which are generally increased in the sleeping period. At the beginning of the lights-off period, HR also rises during cardiac autonomic blockade. The mechanisms of this increase of HR are
unclear but may depend upon behaviorally and sympathetically induced changes in cardiac filling.

Several mechanisms could be responsible for the rise in peripheral resistance during sleep. The most favored explanation is that due to decreased metabolic demand, autoregulatory mechanisms reduce blood flow, especially through skeletal muscle. Alternatively, the diurnal variation in epinephrine, which in rats runs in parallel with the rhythm of HR, may have a role in the rhythm of peripheral resistance via \( \beta_2 \)-adrenergic vasodilatation. The matter becomes even more complicated, because cardiac autonomic blockade attenuated the rise in peripheral resistance at the onset of the inactive period and abolished the relative dark-light difference, especially in SHR (Table 5.1). Thus, under these conditions, the relative changes in TPR are not in phase any more with the dark-light cycle. However, in cardiac autonomic-blocked SHR, peripheral resistance fell during the period when CO was highest, which was at the transition from the light to the dark phase (see Fig. 5.2). The infusion of the drugs may have altered the entrainment to the dark-light cycle and phase advanced the activity pattern of the SHR. Mechanistic studies are needed to resolve to what extent flow and autonomic-dependent mechanisms contribute to control of 24-h peripheral resistance.

**HF oscillations of CO**

Similar to findings in humans, HF oscillations of CO were mainly confined to the respiratory frequency band. As indicated by the normalized transfer function, the relative amplitude of CO was a factor of 7-10 greater than that of blood pressure and not different between SHR and WKY. Furthermore, the phase angle between CO and blood pressure, being close to zero at these frequencies, indicates that the oscillations occur simultaneously. Blockade of the cardiac autonomic nervous system did not alter this. These findings indicate that the respiratory-related oscillations in CO are not related to reflex-induced changes in sympathetic or vagal nerve activity. Thus, these fluctuations are probably induced by a direct mechanical coupling of respiration and CO. The oscillations in CO are then further transmitted into blood pressure and thus certainly not the result of baroreceptor activation.

In the MF, spectral density power of blood pressure, but not of CO, was significantly greater in SHR than in WKY. Similar results were found during cardiac autonomic blockade. These findings suggest that enhanced MF oscillations in the vasculature of SHR may have been responsible for the increased MF power of blood pressure. Indeed, the MF fluctuations in peripheral resistance were higher in SHR than in WKY; however, the difference was not statistically significant. This may be due to an artifact. By calculating peripheral resistance as the quotient of arterial pressure and CO, one transfers the HF oscillations existing in CO to peripheral resistance, where they may not actually exist. Whether the enhanced MF fluctuations in blood pressure in SHR are due to elevated sympathetic nerve activity or exaggerated postsynaptic mechanisms cannot be decided from this study. We favor the latter explanation, because MF oscillations of sympathetic nerve activity do not differ between SHR and WKY.

**LF Oscillations of CO**

Dominant LF oscillations in CO were not found in SHR or WKY either during control conditions or during cardiac autonomic blockade. In all cases, the double loga-
Chapter 5

Rhythmic power spectrum was linear in the frequency band from 0.01 to 0.5 cycles/min. Similar spectra were found for arterial blood pressure and peripheral resistance. These $1/f$ characteristics are typical for systems that are controlled in a nonlinear, chaotic fashion. This means that the system is controlled by multiple processes, acting at overlapping time scales. The slope of the $1/f$ regression line is an index for the complexity of the dynamic process. In the present study, the $1/f$ slopes of the time series of CO, blood pressure, and peripheral resistance did not differ either among themselves or between SHR and WKY. This suggests that, in this LF range, the complexity of the regulation of the systemic hemodynamics is comparable and not dependent on the pressure level. A similar conclusion was reached by Holstein-Rathlou et al. using telemetry to record blood pressure in normotensive and two-kidney, one-clip hypertensive Sprague-Dawley rats as well as SHR. In addition, the absolute values of the slope of the logarithmic regression line between frequency and spectral power of blood pressure were very similar to those reported by that group. During cardiac autonomic blockade, the slope of the $1/f$ time series of peripheral resistance rose significantly in both WKY and SHR. This may be a reflection of the increased impact of baroreceptor-mediated changes on vascular resistance when the autonomic input to the heart is blocked. Evidence is accumulating that the baroreflex buffers low-frequency variations in arterial pressure. In SHR after cardiac autonomic blockade, the slope of the $1/f$ spectrum of arterial pressure increased slightly but not significantly. This finding is consistent with a recent study in sinoaortic-denervated dogs, in which the $1/f$ slope of the blood pressure spectrum was higher than in nondenervated animals. Cardiac autonomic blockade decreased total power of CO, but the $1/f$ slope did not change. This suggests that factors other than the cardiac autonomic nervous system are an important determinant of the LF dynamics of CO.

In contrast to what was found at the high frequencies, the normalized transfer gain for CO and arterial pressure was close to 1 over the whole frequency range. Thus, at these low frequencies, fluctuations of CO and arterial pressure were relatively of the same magnitude. During cardiac autonomic blockade, the relative fluctuations of CO fell whereas those of blood pressure remained unaltered. This reduction was largely due to decreased variability of HR (spectra not shown). The LF oscillations of CO and arterial pressure occurred simultaneously whether or not the cardiac autonomic nervous system was blocked. To our knowledge, dynamic characteristics of cardiac output have been investigated over periods up to 9 h only in one study in dogs. In this report, dominant oscillations with periods near 1.5 h were found in CO and arterial pressure, but the oscillator was not identified. In the present study in rats, such slow rhythms were never observed. However, in both species at these low frequencies, the phase angle between CO and blood pressure was not different from zero. Thus LF fluctuations of CO most likely drive those of arterial pressure.

### 24-h frequency distributions of systemic hemodynamics

The combined frequency distributions of MAP and CO can be regarded as plots of normalized pressure versus normalized flow using the spontaneous variations of these two parameters. In such a plot, complete autoregulation of systemic blood flow would be found when the flow variation remained close to the average value over the whole pressure range. In our plots, however, the relative variation of CO was always greater than that of arterial pressure. This indicates that, over 24-h periods in rats,
Circadian control of cardiac output

arterial pressure was controlled more tightly than CO. In the majority of the rats, the contour plots of MAP and CO and MAP and TPR exhibited a single peak. This could mean that there is one single set point for the combination of these parameters. Alternatively, one may argue that these parameters are fairly unrelated throughout the day and that the contour plot is the result of two gaussian distributions. The contour plots of CO and TPR showed a clear negative correlation, as would have been expected from the diurnal pattern. The width of the distribution was relatively narrow, especially compared with the width of the histograms relating the other combinations. This is another indication that control of pressure is relatively tighter than control of CO or peripheral resistance over 24 h. During cardiac autonomic blockade, coefficients of variation of arterial pressure variations became significantly greater, whereas those of CO decreased slightly. Thus CO fluctuations do stabilize to a certain extent those in arterial pressure. However, in all cases, it was never observed that fluctuations in CO were smaller than those in arterial pressure. This finding seems in contrast with theories that long-term control over CO prevails over control of blood pressure. Over long periods of time, CO is linked to metabolic demand for oxygen and should be independent of neural control. However, in the present study, average CO fell during autonomic blockade, and coefficients of variation were not lower than those of arterial pressure. Possibly large 24-h differences in the metabolic needs of the different tissues may underlie the considerable 24-h labile behavior of CO. Clearly, the present data indicate that 24 h is not enough time for autoregulation of CO to occur.

The existence of multiple states in the contour plots of MAP and PI confirms our earlier findings that multiple preference states exist of combinations of blood pressure and PI. The absence of multiple states in the plots relating the mutual interactions of CO, arterial pressure, and peripheral resistance and the absence of multiple states in the plots relating CO and SV (data not shown) indicate that these preferred combinations are introduced by the variation of PI only. These preference states of PI are not caused by variations in autonomic tone to the heart. In contrast, after cardiac autonomic blockade the number of such preference states in PI even increased. Thus autonomic activity seems to mask the existence of such preferred states. The origin of the preferred states of PI may be related to an intrinsic property of the heart or of the cardiac pacemaker cells in reaction to the opposed work load over 24 h. The preference for a limited number of different frequencies could be a reflection of the existence of subpopulations of pacemaker cells within the sinus node, which have the highest firing frequency, depending on factors such as atrial stretch or humoral factors. On the other hand, it may be energetically more effective to switch between such states than to have a continuous range of HR. A switch between such states is typical for chaotic control systems. Thus the rat heart works in gears rather than on the basis of an automatic transmission.

In summary, using different approaches in examining beat-to-beat recordings of systemic hemodynamics, we found that, over 24-h periods in WKY and SHR, blood pressure seems to be controlled within a more narrow range than CO. Short-term oscillations of CO were, even in the absence of cardiac autonomic tone, coupled to the respiratory cycle. The diurnal variation of CO was reduced but not abolished by blockade of the cardiac autonomic nervous system. This suggests that factors other than the autonomic nervous system control the 24-h. fluctuations of CO. A remarkable finding
Chapter 5

was that the rat heart seems to beat at a limited number of preferential HR. The origin of this property of the heart will need further elucidation.

References


Chapter 6
Time-Dependent Efficacy of Antihypertensive Agents in Spontaneously Hypertensive Rats

Ben J. A. Janssen
Jan Oosting
Caroline M. Tyssen,
Harrie A. J. Struyker-Boudier

Published in Chronobiology International 1993 ; 10, 420-434
Chapter 6

Abstract

The efficacy of antihypertensive agents was compared when given at different time points in the circadian rhythm. Spontaneously hypertensive rats (SHRs) were kept on a 12/12-h cycle with lights on/off at 07:00/19:00 h. A computerized system was used to measure intraarterial blood pressure and heart rate continuously. Agents or vehicle were intravenously injected at two time points. One at the beginning of the sleeping period, at which low efficacy was expected (T = 10), and one at T = 16, which is 3 h before the circadian peaks in blood pressure (BP) and heart rate (HR), aimed at reducing the rise in BP and HR at awakening. The hypotensive effect of propranolol, metoprolol, labetalol, prazosin, clonidine, and rilmenidine was greater when injected at T = 16 than at T = 10 (p < 0.05 for propranolol, metoprolol, and rilmenidine). In contrast, the renal vasodilators captopril and tertatolol were more potent after injection at T = 10. Fentidine was equally effective at both time points. Thus, the effects of antihypertensive agents are related to the phase of the circadian rhythm. The data on the sympatholytic agents in general and β-blockers and centrally acting agents in particular support antihypertensive regimens with timed administrations.
Introduction

The concept of chronopharmacological treatment of hypertension is not new. Its application has regained interest after epidemiological data showed that cardiovascular pathophysiological events (such as sudden cardiac death, myocardial infarction, and ischemia) exhibit circadian rhythmicity too, showing a peak incidence during the early morning hours when blood pressure increases rapidly. Furthermore, meta-analyses of major clinical trials on the benefits of antihypertensive therapy revealed that the reductions in numbers of deaths due to cerebrovascular diseases is about twice that due to cardiovascular diseases. It has been suggested that one of the reasons for this disparity may be that patients are vulnerable to unsuppressed peak morning blood pressure. On the basis of these studies, one may assume that most benefit could be gained by optimizing antihypertensive treatment during the early morning hours when most life-threatening cardiovascular events take place. On the other hand, because most antihypertensive agents lower blood pressure throughout the 24-h period without changing the circadian pattern, improving benefit may depend more on the tonic level of blood pressure or on the choice of the drug. Prospective studies, however, are necessary for proving the beneficial effects of both therapies.

In the present study in spontaneously hypertensive rats (SHRs) we compared the antihypertensive efficacy of agents when administered at different time points in the circadian rhythm. Agents of different classes of antihypertensives were used; sympatholytic agents were investigated in particular. In a previous study in SHR we have shown that these latter agents were more effective in modulating circadian rhythmicity of blood pressure (BP) and heart rate (HR) than nonsympatholytic antihypertensives. Two time points were chosen: one at which low efficacy was expected (i.e., during the beginning of the sleeping phase of the rat) and one at which high efficacy was expected for blocking the spontaneous rise in BP and HR during the awakening period. To avoid chronopharmacokinetic differences in absorption, we administered agents as intravenous bolus injections.

Materials and Methods

Animals

Male adult SHRs (14-16 weeks old) were obtained from the inbred colonies of rats of the central animal facilities of our university. Originally, these rats were derived from the Okamoto-Wistar strains. Before the experiments, rats were housed in groups of four. Rats were maintained on a 12/12-h light/dark cycle (lights on from 07:00-19:00 h, >200 lux) in a quiet room at 21 ± 2EC. Food and water were available ad libitum.

Surgery

Arterial catheters were constructed from a 5-cm PE-10 tubing, heat-sealed to a 80-cm PE-60 tubing. Venous catheters were made from an 80-cm piece of PE-10 tubing, heat-sealed to a 5-cm piece of PE-20 tubing. The PE-10 section of tubing was bent into a J-shape 4 cm from its tip to provide proper angulation of the catheters during implantation. Under aseptic conditions, surgery was performed during light ether anesthesia.
Chapter 6

A longitudinal incision was made in the groin of the leg as well as on the back of the rat 5 cm above the tail. The arterial and venous catheters were implanted via the femoral vessels. The cannulae were advanced 4 cm into these vessels so that the tips were ~1 cm above the bifurcation. The catheters were secured to the leg muscle and tunneled to the back of the animal. The arterial and venous catheters were guided through a stainless steel spring (60-70 cm long) that was fixed to a piece of silicone rubber, which was tightly sutured to the back of the rat with the aid of mersilene gauze. In this way the catheters were protected from damage. The wounds were closed with fine intracutaneous stitches, and the rats were given 150 mg of chloramphenicol per kg. The catheters were filled with heparinized (5 U/ml) saline and together with the spring connected to the equipment as described below.

Measurements

During the experiments, rats were housed individually in their home cages. The arterial catheter was connected via a hydraulic swivel (model 375/20, Instech Labs, Plymouth Meeting, PA, U.S.A.) to a low-volume displacement pressure transducer (micro-Switch, model 156PC156W1, Honeywell, Amsterdam, The Netherlands). A constant perfusion of the arterial line with heparinized saline (30 IU/ml; 2.4 ml/24 h) was maintained by a syringe pump (Razel model A, Razel Scientific Instruments, Stamford, CT, U.S.A.) to prevent clotting in the catheter. The arterial pressure transducer was connected to an amplifier that delivered a high-level voltage signal to an analog-to-digital converter board (model 2814, Data Translation, CN Rood, Rijswijk, The Netherlands) housed in an IBM AT-compatible microcomputer. Electrical drift in the arterial pressure monitoring system was acceptably low and smaller than 1 mmHg/week as revealed by periodic calibrations. The system allowed simultaneous recording of eight rats. The arterial pressure signal was sampled continuously at ~60 samples/beat. Software was developed for calculating beat-by-beat mean arterial pressure (MAP) and pulse interval (PI). These beat-to-beat values for MAP and HR (60,000/PI) were averaged each min. Hourly values for MAP and HR were calculated by averaging 60 1-min values and stored on disk.

Protocols

Experiments were started after rats had recovered at least 72 h from surgery. From then on reproducible circadian rhythms in BP and HR can be obtained. Drugs were given as slow bolus injections (0.1-0.3 ml) via the venous catheter that was extended outside the cage, followed by 0.25 ml saline for washing the cannula. In this way injections can be given without disturbing the rats. Injections were given at two different time points in the circadian cycle. Vehicle (saline) or drug solutions were injected either at 10:00-10:15 h (T = 10) or at 16:00-16:15 h (T = 16), and effects on MAP and HR were studied for 24 h. At least 48 h were allowed to pass before another injection was given. Drugs were given in random order. The time of injection was also randomized. The following antihypertensive agents were used: metoprolol (1.5 mg/kg), propranolol (5 mg/kg), prazosin (0.6 mg/kg), and labetalol (10 mg/kg), respectively, as a β-1 selective, a β-nonsellective, an α-1 selective, and combined β- and α- adrenoceptor antagonist; the centrally acting agents clonidine (10 µg/kg) and rilmenidine (1 mg/kg) [rilmenidine was chosen because this agent is less sedative than clonidine]1; the angiotensin I converting enzyme inhibitor captopril (10 mg/kg); tertatolol (0.5 mg/kg) as a β-blocker
with 5-HT1ₐ antagonistic properties; and the calcium antagonist felodipine (0.3 mg/kg).
As an effective dose, onethird was taken of the amount used in a previous experiment
while studying the efficacy of continuous infusions¹⁰. Agents were dissolved in saline
or, in the case of propranolol, prazosin, and captopril, in water.

**Calculations and Statistics**

Because 24-h MAP and HR rhythms vary from day to day, for each individual rat
a 24-h reference curve was calculated by averaging MAP and HR over at least three, but
in most cases four, different 24-h periods during which only saline was injected. The
effects of drugs on MAP and HR are then related to this reference curve. The duration
of the effect of each single injection could not be determined accurately because of the
individual spontaneous variability in MAP and HR. Therefore, the averaged (over all
rats) time curves of MAP and HR were used in determining the mean duration of an
effect (TDurMap and TDurHR respectively). These averaged time curves as well as the
averaged reference curves were subjected to a cumulative summation technique, to
construct a so-called “cumumplot,” from which time points of change (end of effect) can
be identified¹¹.

Then for each single injection, a summation was made of the differences in MAP
and HR with the reference curve over TDurMap and TDurHR. These latter values of
summed differences in MAP and HR can be regarded as a measure of the pharmacodynamic response and were defined as \( \Delta \text{Map} \) and \( \Delta \text{HR} \). They are visualized as the
shaded area in Figs. 1-9. Furthermore for each injection the maximal effect \( (\Delta \text{Map}_{\text{max}} \text{ and } \Delta \text{HR}_{\text{max}} \), maximal difference with the reference curve) was determined. As a last
parameter of drug efficacy, the difference (with the reference curve) in the summated
rate pressure product (RPP = MAP * HR) over TDurMap was calculated (\( \Delta \text{RPP} \)). Then,
the above named parameters were averaged per agent and per injection time.

For each agent, differences between these parameters at injection time T = 10 or
T = 16 were compared with a paired t-test in the event paired recordings could be
completed for all rats of one group. Otherwise an unpaired t-test was applied. Because
Table 6.1. Comparison of the efficacy of antihypertensive agents.

<table>
<thead>
<tr>
<th></th>
<th>T = 10</th>
<th>T = 16</th>
<th>ΔMAPmax</th>
<th>ΔHRmax</th>
<th>ΔMAPmean</th>
<th>ΔHRmean</th>
<th>ΔSHRmean</th>
<th>ΔRPP%</th>
<th>t-test</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Dose</strong> mg/kg</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n at T = 10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>n at T = 16</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>ΔMAPmax</td>
<td>-6.3</td>
<td>-6.2</td>
<td>-11.6</td>
<td>-5.9</td>
<td>-8.6</td>
<td>-7.7</td>
<td>-5.6</td>
<td>-3.3</td>
<td>-2.5</td>
</tr>
<tr>
<td>ΔHRmax</td>
<td>-5.9</td>
<td>-4.4</td>
<td>-2.8</td>
<td>-1.1</td>
<td>-1.6</td>
<td>-1.8</td>
<td>-1.8</td>
<td>-1.8</td>
<td>-1.7</td>
</tr>
<tr>
<td>ΔMAPmean</td>
<td>2.9</td>
<td>3.6</td>
<td>5.1</td>
<td>3.5</td>
<td>3.3</td>
<td>3.5</td>
<td>3.3</td>
<td>3.2</td>
<td>3.3</td>
</tr>
<tr>
<td>ΔHRmean</td>
<td>8.2</td>
<td>7.7</td>
<td>6.8</td>
<td>7.2</td>
<td>7.5</td>
<td>7.7</td>
<td>7.5</td>
<td>7.5</td>
<td>7.5</td>
</tr>
<tr>
<td>ΔSHRmean</td>
<td>1.1</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
</tr>
<tr>
<td>ΔRPP%</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>unpaired</td>
<td>unpaired</td>
<td>unpaired</td>
<td>unpaired</td>
<td>unpaired</td>
<td>unpaired</td>
<td>unpaired</td>
<td>unpaired</td>
<td></td>
</tr>
</tbody>
</table>

Agents were injected at T = 10 or T = 16. Effects are expressed by maximal and summed changes in mean arterial pressure (ΔMAPmax, ΔMAP) and heart rate (ΔHRmax, ΔHR) and differences in rate pressure product (ΔRPP) over the duration of the effect. *P < 0.05 significantly different between effects at T = 10 and T = 16.

no dose response curves were constructed, differences between agents were not subjected to quantitative analyses, but described only qualitatively. Data are presented as means ± SE unless indicated otherwise. Differences were regarded significant at P < 0.05.

Results

As depicted by the reference curves in Figs. 6.1-6.9, MAP and HR vary considerably in SHRs throughout a 24-h period. However, there is a consistent pattern. MAP and HR are low during the lights-on period with a small peak at ~ 10:00 A.M. Then, from ~ 16:00 h, MAP and HR start to rise, exhibiting a first peak at 20:00 h followed by two other peaks at ~ 00:00 and 6:00 A.M. during the lights-off period. Maximal MAP and HR values were usually found during the third and first peak of the dark period, respectively. Figures 6.1-6.9 compare for each agent the effects of a single injection on MAP and HR when given either at T = 10 or T = 16. Table 6.1 summarizes the results for the deducted parameters such as maximal and summed difference in MAP and HR as well as differences in RPP caused by the injections. The duration of the averaged response to each drug is indicated in the figures by the shaded area.

The antihypertensive effect of metoprolol (Fig. 6.1) was significantly greater when
the agent was injected at T = 16 than at T = 10. At T = 10 there was even a slight increase in HR, whereas at T = 16 HR was lower than the reference curve, but increasing in a similar way as the reference curve.

When injected at T = 10 propranolol (Fig. 6.2) first increased MAP for 2 h and then decreased it for 4 h, accompanied by a small increase in HR. At T = 16 the initial increase in MAP was smaller than after T = 10 and was followed by a greater and much longer-lasting reduction in MAP. Reductions in HR after injection of propranolol at T = 16 were small but significantly different from those at T = 10. For both propranolol and metoprolol the reduction in ΔRPP was significantly greater after injection at T = 16 than T = 10.

Prazosin (Fig. 6.3) reduced MAP effectively when injected either at T = 10 or T = 16. ΔMAP, ΔMap, and ΔRPP were greater at T = 16 than at T = 10, although differences were not statistically significant. The parameters describing the degree of tachycardia following administration of prazosin were greater at T = 10 than at T = 16.
These differences, however, also did not reach significance.

After injection of labetalol at $T = 16$, $\Delta \text{Map}$ and $\Delta \text{MAP}_{\text{max}}$ were greater than at $T = 16$, but differences were not statistically significant. At both injection times HR increased following administration of labetalol. Maximal changes in HR were not different, but the degree of tachycardia ($\Sigma \Delta \text{HR}$) was significantly greater at $T = 10$ than at $T = 16$.

The BP lowering effect of clonidine (Fig. 6.5) was greater when injected at $T = 16$ than at $T = 10$. The differences in $\Sigma \Delta \text{Map}$ (-36 ± 12 vs. -51 ± 12 mm Hg) were almost significant ($p < 0.08$). The decreases in HR and $\Delta \text{RPP}$ following clonidine administration were similar at $T = 10$ and $T = 16$.

Rilmenidine (Fig. 6.6) decreased MAP and HR effectively in SHRs. Maximal reductions in MAP and HR that were achieved after injection at $T = 10$ or $T = 16$ were comparable. The summation of MAP and HR reductions as well as the reduction in
In contrast to the above-mentioned sympatholytic agents, captopril (Fig. 6.7) was longer effective in lowering MAP when given at $T = 10$ rather than at $T = 16$. However, values of $\Sigma\Delta$Map at $T = 10$ and $T = 16$ were not significantly different. In both cases the effects on MAP were accompanied by a small degree of tachycardia.

A similar pattern was found when tertatolol (Fig. 6.8) was given. Reductions in $\Sigma\Delta$Map were approximately three times greater when given at $T = 10$ than at $T = 16$. However, the variability in this response was quite large, and differences between mean values were not statistically significant. Remarkably, tertatolol was without effect on HR.

The antihypertensive effects of felodipine (Fig. 6.9) were similar when given at either $T = 10$ or $T = 16$, although the duration of the response seems to be greater after injection at $T = 10$. At both times, the agent caused a profound reduction in MAP.
associated with tachycardia. After the maximum effect was reached, MAP returned rapidly to control levels.

**Discussion**

The present study shows that the cardiovascular effects of a number of antihypertensive agents in SHRs are dependent on the time of administration in the 24-h cycle. Sympatholytic agents, in particular, were consistently more effective when administered at the beginning of the rise in BP at awakening than during the first hours of the sleeping period.

Both, circadian phase-dependent changes in pharmacokinetics and pharmacodynamics may underlie the temporal differences in efficacy of the agents. Chronokinetic differences in reabsorption were avoided by giving intravenous bolus injections. How-
ever, temporal differences in binding capacity of plasma proteins, metabolic clearance rate, or excretion may influence the availability of agents. Yet, the present data on the β-blockers propranolol and metoprolol suggest that chronokinetic factors are of minor importance as compared with chronopharmacodynamic factors. Whereas the antihypertensive efficacies of propranolol and metoprolol were greater when injected at T = 16 than at T = 10, the elimination half-lives of these β-blockers in plasma and organs of rats are known to be shorter during the dark period than during the light period. Similar conclusions were reached when the pharmacokinetics and cardiovascular effects of propranolol in healthy subjects were compared. It has been suggested that increased liver blood flow and increased metabolic activity during the active period, on the one hand, and increased susceptibility due to enhanced sympathetic activity, on the other hand, underlie the discrepancy between pharmacokinetic and pharmacodynamic behavior of these β-blockers.

Although no dose-response studies were performed and no quantitative comparison of the differences in efficacies of the agents can be made, certain aspects of the present findings require specific discussion. The most clear-cut circadian temporal differences in efficacy were found with the β-blockers metoprolol and propranolol. The antihypertensive response was much greater and longer lasting when these agents were injected at T = 16 than at T = 10. In contrast to that of metoprolol, the antihypertensive effect of propranolol at T = 10 and T = 16 was preceded by a 2-h and a 1-h lasting increase in blood pressure, respectively, which is probably caused by baroreceptor-mediated peripheral vasoconstriction.

The injection of the β-blockers at T = 16 did not cause a profound bradycardia. The maximal average reduction in HR in 1 h was only 11 beats/min for both the β-blockers. The rate of rise in HR during that period was also not different from the nontreated situation. This finding suggests that the increase in HR during the awakening rise in rats is probably more the result of a fall in parasympathetic activity rather than sympathetic activation. When these β-blockers were injected during the first hours of the sleeping period, during which basal HR is very low and vagal activity prevails, even small, but reproducible, average increases in HR were found. Since both agents lack intrinsic sympathicomimetic activity, the mechanism of this response is unclear. However, we speculate that a β-blocker-induced fall in stroke volume may underlie this small increase in HR. On the other hand, injection of propranolol at T = 10 may have unmasked a vagal cardioacceleratory system in these rats.

The effects caused by the α-blocker prazosin and labetalol, an agent with combined α- and β-blocking properties were comparable and for a major part determined by baroreceptor reflex activation. Labetalol was used since in human studies this agent was most effective in reducing the morning rise in BP. In SHRs prazosin and labetalol caused a pronounced fall in MAP, and the hypotensive response was greater when the agent was injected at T = 16 than at T = 10. Differences in these effects at T = 10 and T = 16 were not statistically significant and may be due to the fact that the administered dose was quite large for both agents. After injection of labetalol, the amount of reflex tachycardia was greater at T = 10 than at T = 16. Not only maximal changes in HR were larger, but the duration of the reflex tachycardia was also enhanced after injection at T = 10. Differences in the central processing of baroreceptor reflex arch as well as differences in pharmacokinetics may explain this finding. Baroreceptor sensitivity has been
Chapter 6

reported to be enhanced during sleeping hours \(^9\), which is corroborated by the fact that BP lability indexes are generally lower during these hours.\(^{10,11}\) On the other hand the rate of elimination of labetalol may have been increased due to the enhanced metabolic activity during the dark hours. Further studies should clarify whether and how prazosin and labetalol can be administered for an optimal chronopharmacological profile.

The cardiovascular effects of the centrally acting agents clonidine and rilménidene were also more pronounced at \(T = 16\) than at \(T = 10\). These data are in agreement with those of Sannajust et al.\(^{29}\), who showed that the BP lowering effect of these agents in SHRs is related to the preexisting sympathetic tone. In addition, the hypotensive effect of clonidine and rilménidene at \(T = 16\) may also be related to their inhibitory effect on vasopressin and adrenocorticotropic hormone (ACTH) release\(^{21}\). These latter effects may be more pronounced at \(T = 16\), since this is the time point at which plasma corticosteroid levels are supposed to peak\(^{22}\). In contrast with the \(\beta\)-blockers, the effects of clonidine and rilménidene were accompanied by bradycardia, even when injected at \(T = 10\), when sympathetic activity is supposed to be minimal. This reduction in HR may be explained by baroreceptor loading caused by an initial rise in pressure as a result of vascular \(\alpha\)-adrenoceptor stimulation. The \(\beta\)-blockers, in contrast, lead to baroreceptor unloading. Thus, the effects of clonidine and rilménidene seem to be governed by vagal activation as well as sympathetic inhibition.

The temporally dependent cardiovascular effects of the angiotensin-converting enzyme (ACE) inhibitor captopril were opposite to those found after injection with the above-mentioned sympatholytic drugs. Although \(\Sigma\Delta\text{MAP}\) at \(T = 10\) was twice the value found at \(T = 16\), these differences in efficacy did not reach significance because of considerable intraanimal variation. Part of this variability may be explained by the fact, as was found in humans\(^{33}\), that plasma renin activity (PRA) shows a burst-like mode of appearance over 24 h. Especially during the sleeping period, oscillations in the behavior of PRA coincides with episodes of rapid eye movement (REM) and non-REM\(^{34}\). Remarkably, the duration of the BP lowering effect of captopril when injected at \(T = 10\) was much longer than that observed after injection at \(T = 16\). Similar findings were observed after injection of tertatolol, a \(\beta\)-blocker with \(5\text{-HT}\text{1}_{\text{A}}\) receptor-blocking properties\(^{35}\). The in vivo cardiovascular effects of captopril\(^{36}\) and tertatolol\(^{37}\) have been reported to last considerably longer than can be expected on the basis of their half-lives. The underlying hemodynamic mechanisms of this prolonged cardiovascular activity may depend on the specific renal vasodilatory properties of captopril and tertatolol\(^{38,29}\). Thus, especially during the sleeping period, when urine production and glomerular filtration rate are lower than during the activity period\(^{38}\), the kidney may be, as has been shown for furosemide\(^{31}\) and a preparation of captopril and hydrochlorothiazide\(^{32}\), highly sensitive to renal vasodilators. Further studies are necessary for supporting this hypothesis, which may lead to a timed regimen of administration of ACE inhibitors.

The antihypertensive efficacy of felodipine in SHRs was not dependent on circadian phase. Recently, similar results were obtained in SHRs for amloidipine by Lemmer et al.\(^{33}\). On the other hand, this group reported that the effects of amloidipine in normotensive rats were dependent on circadian phase\(^{34}\). Thus, strain differences seem to influence the circadian efficacy of calcium antagonists.

In summary, the cardiovascular effects of antihypertensive agents in SHRs are dependent on their time of administration. Especially, \(\beta\)-adrenergic blockers and cen-
Chronopharmacology of antihypertensive agents

tially active agents showed greater antihypertensive efficacy when given at a time point to attenuate the early morning rise (in rats, early night) in BP. This finding suggests that agents that lower BP, among other mechanisms, by a reduction of cardiac output, may be suitable tools for the chronopharmacological treatment of hypertension.

References


17. Struyker Boudier HA, Smits JFM, van Essen H: The role of the baroreceptor reflex
Chapter 6


Chapter 7
Circadian-phase dependent pharmacodynamics of angiotensin converting enzyme inhibitors in spontaneously hypertensive rats

Jan Oosting
Harry A. Struijker-Boudier
Ben J. Janssen

Submitted for publication
Chapter 7

Abstract

Objective To study the antihypertensive efficacy of drugs that inhibit the renin angiotensin system when administered at different time points of the day.

Methods Adult spontaneously hypertensive rats, were treated on 5 consecutive days with infusions of the ACE-inhibitors captopril (3, 10, and 30 mg.kg⁻¹.6 h⁻¹), enalaprilat (0.3 mg.kg⁻¹.6 h⁻¹), lisinopril (0.1 mg.kg⁻¹.6 h⁻¹), or the AT₁-receptor antagonist losartan (10 mg.kg⁻¹.6 h⁻¹), either during the transition from the lights-off to the lights-on, resting period (evening dose), or during the transition from the lights-on to the lights-off, active period (morning dose).

Results The effects of all agents on the circadian blood pressure pattern were comparable. In general, the average reduction of blood pressure over 24 hours was not different following a morning or evening dose. However, the blood pressure patterns were markedly different during specific periods of the day for the two treatment regimens. The difference in blood pressure between the lights-off and lights-on period was smaller following the morning than the evening treatment regimen. Reflex tachycardia was more pronounced when drugs were given as the evening dose, whereas the rate pressure product, an index of cardiac oxygen consumption, was more decreased following the morning dose.

Conclusions In spontaneously hypertensive rats, circadian rhythms of blood pressure can be modulated by timed administration of agents that inhibit the renin angiotensin system. Especially when given during the transition from the resting to active period of the day, these agents reduce effectively the surge in blood pressure normally observed in this period.
Introduction

Morbid events related to hypertension, such as myocardial infarction, angina pectoris, and nonembolic strokes strike more often in the early morning hours than at other time points of the day\textsuperscript{1-4}. This has raised the question whether protection against such events can be enhanced by targeting the anti-hypertensive therapy towards this early morning period, which is characterized by a sharp rise in blood pressure and heart rate\textsuperscript{5,4}. Prospective clinical studies in support of this hypothesis are not available. Nowadays one special delivery system has been developed for the calcium antagonist verapamil. The drug preparation is designed to control the onset and to extend the release of the agent to achieve maximal levels of the drug during the early morning surge in blood pressure\textsuperscript{6}.

For angiotensin converting enzyme (ACE) inhibitors, such special formulations do not exist. Obviously, this can be explained by the fact that recently developed ACE inhibitors have a long pharmacological half-life and are applicable as a once a day preparation aimed at controlling blood pressure over the whole 24-h period. However, when therapy once a day is used, the drugs are usually ingested in the morning. Hence, one may expect that the lowest plasma levels and the least pharmacological effect will occur during the last hours of the dosing interval, which coincides with the early morning blood pressure rise. Thus, there may be insufficient control during the time at which hypertensive patients are at greatest risk for the development of cardiovascular events. Indeed, studies on once a day B-blockers\textsuperscript{6} and ACE inhibitors\textsuperscript{7-10} have indicated that blood pressure control may not be adequate during the last hours of the 24-h dosing interval, particularly when the drug is ingested as a morning dose.

To obtain a significant fall in blood pressure at the end of the dosing interval, the United States Food and Drug Administration (FDA) introduced the concept of the trough-to-peak ratio. New antihypertensive agents should meet the requirement that 50-65% of their maximal blood pressure lowering effect (peak) persists until the next dose (trough)\textsuperscript{11-12}. Dose titration of antihypertensives is complicated, also when drugs have a relatively linear relation between their pharmacokinetic and pharmacodynamic behaviour\textsuperscript{13}. This process may become even more complex when a direct relation between these parameters is lacking. This applies particularly for many ACE inhibitors, since their antihypertensive effect depends partly on the amount of inhibition of tissue ACE\textsuperscript{14,15} and does not coincide with the time when peak plasma levels are achieved\textsuperscript{6,16}.

Many mechanisms have been brought forward to explain the pharmacodynamic behaviour of ACE inhibitors. Relevant to this study, is the circadian variation in the components of the renin-angiotensin system. In normal and hypertensive humans\textsuperscript{17,18} and in animals\textsuperscript{19,20} plasma renin levels are low at the beginning of the sleeping period but increase steadily to be highest at the time of awakening. However, little to none information is available about the circadian variability in plasma and tissue of other components of this humoral system\textsuperscript{21}. Despite the lack of such basic data a few recent studies in patients have demonstrated that the timing of dosing is important in the 24-h protection by ACE inhibitors\textsuperscript{5,10}. The results of these studies are not very consistent, which may be one of the reasons why a chronotherapeutic approach of hypertension has received little attention.
Chapter 7

To generate more conclusive information about the time-dependent efficacy of ACE-inhibitors in the treatment of hypertension, we investigated the circadian-phase dependent efficacy of several ACE inhibitors in spontaneously hypertensive rats (SHR). This animal model was chosen because the circadian blood pressure rhythm in this species is comparable to the one in essential hypertensive patients. One should, however, account for a 12-h phase difference in the haemodynamic patterns in these nocturnally active animals. In SHR, blood pressure and heart rate rise slowly during the last two hours of the lights-on period and increase rapidly when the rats become fully active when the lights are off. At the beginning of the lights-on period blood pressure and heart rate fall steeply. A second reason to use this model was that, comparably to findings in patients, the circadian pattern of blood pressure is preserved, when antihypertensive agents are continuously infused. For instance, during a constant intravenous infusion with captopril, we found that the circadian variation in blood pressure was not altered despite a 20-30 mmHg reduction in arterial pressure. Assuming that under these circumstances steady state inhibition of ACE was achieved, this suggests that the pharmacodynamic reaction to an ACE inhibitor is more important than its pharmacokinetic profile. Lastly, in two studies in SHR, the antihypertensive effect following single bolus injections of captopril and enalaprilat were circadian phase-dependent too.

In the present study, we have compared the antihypertensive efficacy and trough-to-peak ratio's of the ACE-inhibitors, captopril, enalaprilat and lisinopril, and the AT,-receptor antagonist losartan when infused intravenously in SHR during the transition from the dark to light period or during the transition from the light to dark period. All agents were infused at either time period repeatedly for a period of five consecutive days. We chose to administer the agents intravenously to minimize acute counter regulatory effects associated with bolus injections as well as to obtain comparable bioavailability at both periods of the day. When agents are administered orally, circadian-phase dependent differences in absorption may determine importantly the effect. Captopril was used as a reference drug at three different dosages. To investigate whether central nervous effects of ACE-inhibition could be involved we compared the time-related effects of captopril to those of enalaprilat, which does not cross the blood brain barrier as readily as captopril. The ACE-inhibitor lisinopril was compared with captopril to estimate the influence of its longer half life on the time-related effects on the circadian blood pressure patterns. Lastly, losartan, was used to discriminate between effects of ACE inhibition and AT,-receptor blockade.

Methods

Animals

Adult SHR rats weighing 280-300 g were used. The rats were obtained from the breeding colonies of the central Animal Facilities of the Universiteit Maastricht. Experiments were performed according to the institutional guidelines, and were approved by the institutional animal ethics committee. After surgery, the animals were housed individually in cages, and were allowed normal rat chow and drinking water ad libitum. The animals were kept on a 12-hour lights-on/12-hour lights-off circadian rhythm in a climati-
Figure 7.1. Time line of the experimental protocol. The graph shows 1 week of recording: 5 days of either saline or drug infusion (D1-D5) and 2 days of recovery (R1 and R2). The synchronization in time with the lights-off (black bars) and lights-on (white bars) period is given by the light-dark row. The bars in the row of Morning/Evening indicate the actual 6 hours periods at which the rats received saline or the drug as a morning or evening dose, respectively.

Sedation was performed in a room with constant temperature (22°C) and humidity (70%). Before the measurements, rats were accustomed to the light-dark cycle for at least one week.

Surgery

All rats were instrumented with an arterial catheter for measuring blood pressure and a venous catheter for administration of drugs as described in detail previously. In short, the catheters were implanted via the femoral vessels and then tunnelled to the lower back of the animal under pentobarbital anaesthesia. The catheters were exteriorized through a 40 cm long steel spring which was fixed to the back muscles with a piece of silicone rubber. The venous catheter was filled with a heparinised saline solution (5 U/ml). The steel spring was led to the outside of the cage, where the arterial catheter was connected via a hydraulic swivel (Model 375/20, Instech Labs, Plymouth Meeting, PA, USA) to a low-volume displacement pressure transducer (micro-switch, model 156PC 156WL, Honeywell, Inc, Amsterdam, the Netherlands). The arterial cathula was kept patent by a continuous arterial infusion of heparinised saline solution (36 U/ml) at a rate of 2.4 ml per day. This setup allowed us to measure blood pressure continuously and to administer drugs without disturbing the rats, for periods lasting 5 weeks in general.

Data acquisition

After surgery, the animals were allowed at least 3 days of recovery to restore the circadian rhythm of haemodynamics. The measurements were done in the home cage of the animal. The arterial pressure transducer was connected to an amplifier that delivered a high-voltage signal to an analog-to-digital converter board (model 2814, Data Translation, CN Rood, Rijswijk, the Netherlands) mounted in an IBM compatible computer. The blood pressure signal was sampled at 512 Hz. Beat-to-beat values of mean arterial pressure (MAP), heart rate (HR), and rate pressure product (RPP) as an index of oxygen consumption by the heart, were calculated on-line. All data were averaged every half hour and stored on computer disk for later analysis.
Chapter 7

Table 7.1. Baseline values of mean arterial pressure, heart rate and rate pressure product

<table>
<thead>
<tr>
<th>Evening Dose</th>
<th>Morning Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>MAP (mmHg)</td>
</tr>
<tr>
<td>Captopril 3</td>
<td>9</td>
</tr>
<tr>
<td>Captopril 10</td>
<td>8</td>
</tr>
<tr>
<td>Captopril 30</td>
<td>9</td>
</tr>
<tr>
<td>Enalaprilat</td>
<td>8</td>
</tr>
<tr>
<td>Losartan</td>
<td>8</td>
</tr>
<tr>
<td>Lisinopril</td>
<td>6</td>
</tr>
</tbody>
</table>

Values are 24-h averages over the first control day. Values are expressed as means ± SEM. MAP: mean arterial pressure; HR: heart rate; RPP: rate pressure product. * P < 0.05 morning vs evening dose.

Protocols

Each experiment lasted one full week, during which MAP, HR, and RPP were continuously measured. The design of the protocol is summarized in Figure 7.1. One group of SHR was treated on five consecutive days during the transition from the light to dark period. The infusion of the drug was started 2 hours before the lights went off and was maintained for 6 hours. In parallel with the human situation, this period was defined as the morning dose, since the agents were given during the transition from sleep to activity in the rat. In the other group of SHR the agents were infused five days for 6 hours during the transition from the light to dark period, starting 3 hours before lights on. This infusion period was defined as the evening dose. After a five days treatment period, measurements of MAP, HR, and RPP were continued for 2 days to record the effects of withdrawal of the agent. In each animal a time control experiment was performed consisting of a 5 day repeated infusion of 0.9% NaCl at a rate of 2.4 ml/6 hours either during the morning or evening period followed by 2 days of recovery. The volume of the infused fluid (2.4 ml/day) is not likely to influence haemodynamics regarding the fact that adult rats drink about 30 ml of water daily. The order of such a 7 days experiment with an antihypertensive agent or with saline was randomized in each rat.

Using this protocol, the following sets of experiments were conducted. Captopril infusions were performed as a morning and evening dose at 3 mg.kg⁻¹.6 h⁻¹, 10 mg.kg⁻¹.6 h⁻¹, and 30 mg.kg⁻¹.6 h⁻¹. Enalaprilat was infused as a morning and evening dose at 0.3 mg.kg⁻¹.6 h⁻¹. Lisinopril was infused as a morning dose at 0.1 mg.kg⁻¹.6 h⁻¹ given at the morning period. In addition, lisinopril was administered as an equivalent dose dissolved in the drinking water. Losartan was infused as a morning and evening dose at 10 mg.kg⁻¹. If possible multiple agents were infused in one rat. Treatment was however not commenced before the blood pressure level had returned to control values.
Data analysis

Recordings were made for 7 full days. The beginning of day 1 was defined at halfway the lights-off or dark period at which the first infusion was given (Figure 1). For each day of treatment averages of MAP, HR and RPP were calculated every 30 minutes. The antihypertensive efficacy was calculated as the absolute difference of MAP, HR, and RPP in each 30 minute period with the values obtained during the saline infusion in the same rat. The average blood pressure lowering effect over a complete day was calculated as the average value of 48 half-hour differences in each day. To obtain a measure for the duration and stability of the antihypertensive efficacy, three parameters were calculated additionally. As a first parameter we calculated the maximal amplitude of the circadian rhythm of MAP by subtracting the half-hour minimum value in each day from the half-hour maximal value in each day. As a second parameter the MAP difference between the light and dark period was calculated by subtracting the average 12-h value of MAP during the light period from the average 12-h value during the dark period. For each infusion day a trough-to-peak ratio of MAP was calculated as the quotient of the average MAP value in the hour preceding the next infusion (trough) and the maximum reduction of MAP on 3 consecutive half hour values in the 24-h period after the start of the infusion (peak). The within animal variability of this parameter was determined as the coefficient of variation over the trough-to-peak values obtained on days 3, 4 and 5 of treatment. Since the latter was very high, the trough-to-peak value was also computed from group averages of the trough and peak effects in blood pressure.

Drugs

Captopril was purchased from Sigma Chemicals, Axel, The Netherlands. Enalaprilat and lisinopril and losartan were kindly donated by Merck Sharp and Dohme, Haarlem, The Netherlands. All agents were dissolved in saline and frozen in small aliquots before use.
Figure 7.3. Patterns of mean arterial pressure (MAP) and heart rate (HR) during the 5 days of treatment (D1–D5) with captopril 10 mg kg⁻¹ 6 h⁻¹ as an evening dose (left panel) and as a morning dose (right panel). The effects upon withdrawal of captopril are also shown (R1 and R2). Values are group averages obtained during the infusions of saline (thin line) and during captopril (thick line). The number of animals in each group is given in Table 7.1. For reasons of clarity the SEM is not shown. The closed bars denote the lights-off period of the day. The 6 h infusions were given during the periods denoted by the open bars.

Statistics

Statistical evaluation was performed by analysis of variance (ANOVA). Haemodynamic effects occurring over time during saline or drug infusion in each rat were defined as a within factor. Differences between the effects caused by a morning dose and an evening dose as well as differences in effects between agents were compared as between factors. Data obtained during the first two days of treatment were not included into the analysis because of possible run-in effects of the drugs. Only average values found at days 3, 4, and 5 of treatment were included in the evaluation. Differences between drugs were tested by ANOVA followed by a post-hoc Scheffé's T-test. Statistical significance was accepted at P < 0.05.

Results

Baseline haemodynamics

Baseline haemodynamic values of MAP, HR, and RPP are compared between groups of rats in Table 7.1. The baseline values were defined as the average 24-h values of MAP, HR, and RPP during the first day of intravenous saline infusion. The table shows that these values of MAP, HR and RPP were not different between the group of rats which received saline during the morning period and the groups of rats which
received saline during the evening period. Only between the groups of rats which were treated with 3 mg.kg⁻¹, 6 h⁻¹ captopril a slight but significant difference in the baseline values of HR was found.

In all groups, the five-day repeated infusion of 2.4 ml saline solution in 6 hours did not alter the 24-h pattern of blood and heart rate. As shown in Figures 7.2 and 7.3, the circadian profile of MAP and HR remained unaltered for the 7-day control period when saline was infused during the morning or evening period. Stability of the haemodynamics during the week of saline infusions was assessed in all groups by ANOVA with time as a within factor. The analysis revealed that in none of the groups MAP and HR were significantly different between the days of the week.

For better understanding of the 24-h blood pressure patterns during timed antihypertensive treatment it is necessary to describe in detail the 24-h MAP and HR profiles during saline infusion. As shown in Figure 7.2 the 24-h HR pattern shows a bimodal distribution with high values during the dark period, and low values during the light period. Highest values of HR are found during the transition from the light to the dark period, when the rats become active. The 24-h MAP pattern shows a more gradual increase in blood pressure throughout the day, starting in the light period and showing the highest blood pressure at the end of the dark period (Figure 7.2). During the transition from the light to dark period mostly a sharp peak occurs. During antihypertensive treatment, these 24-h patterns of MAP and HR are generally preserved and visible on top of the effects induced by the agents.

**Time-related effects of treatment with captopril**

The time-related effects during the five-day treatment with captopril 10 mg.kg⁻¹, 6 h⁻¹ are summarized in Figure 7.3. In both experiments the antihypertensive properties of captopril showed increasing efficacy over the ensuing days of treatment with only a small reduction in blood pressure occurring during the first infusion, and more than 30 mmHg during the infusion of captopril at day 5. This effect was more pronounced when
Captopril was given as a morning dose than as an evening dose, in which case a full anti-hypertensive effect was obtained at the third day of infusion. As summarized in Figures 7.4 and 7.5, the average reduction in blood pressure over day 3 to 5 of infusion was not different when the 10 mg dose of captopril was infused in the evening or in the morning period. However, the 12-h phase difference between the infusion periods had a pronounced differential effects on the circadian profiles of MAP. As shown in the Figures 7.3 and 7.4, the evening dose of captopril lowered the peak of MAP which normally occurred at the end of the dark period, and when the lights went on, MAP declined further. The morning dose of captopril blunted the rise of blood pressure in the transition from the light to dark period. After cessation of the 6 h infusion with captopril, MAP returned rapidly to levels not far from control. Reflex tachycardia was significantly greater when captopril 10 mg.kg⁻¹ was given as the evening dose than when given as the morning dose (Figure 7.4 and 7.5). Consequently, also the fall in RPP was significantly greater for the morning dose than evening dose of captopril 10 mg (Figure 7.5).

The result of this pharmacodynamic behaviour of captopril is that the 24-h maximal amplitude of MAP significantly increased from about 32 mmHg during the saline infusion to 44 mmHg during the captopril 10 mg.kg⁻¹.6 h⁻¹ infusions (Table 7.2). The increase in circadian amplitude was comparable for the evening and morning dose of captopril 10 mg.kg⁻¹.6 h⁻¹. Secondly, because of the 12-h phase shift between the infusion periods the differences between the averages of MAP during the 12-h dark and 12-
### Table 7.2. Maximal amplitude of the 24-h blood pressure variation

<table>
<thead>
<tr>
<th>Evening Dose</th>
<th>Morning Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n Control (mmHg)</td>
</tr>
<tr>
<td>Captopril 3</td>
<td>9 34 ± 2.9</td>
</tr>
<tr>
<td>Captopril 10</td>
<td>8 31 ± 2.3</td>
</tr>
<tr>
<td>Captopril 30</td>
<td>9 33 ± 2.6</td>
</tr>
<tr>
<td>Enalaprilat</td>
<td>8 32 ± 1.6</td>
</tr>
<tr>
<td>Losartan</td>
<td>8 33 ± 2.1</td>
</tr>
<tr>
<td>Lisinopril</td>
<td>6 37 ± 1.3</td>
</tr>
</tbody>
</table>

Values are the average (over days 3-5) difference between the minimum and maximum 30 min value of MAP occurring during 24 hours. Recordings were made during saline infusions (control) and during treatment. Values are expressed as mean ± SEM. *: lisinopril in drinking water; †: P < 0.05 treatment vs. control; †: P < 0.05 morning vs. evening dose.

### Table 7.3. Blood pressure difference between 12-h lights-off and 12-h lights-on period.

<table>
<thead>
<tr>
<th>Evening Dose</th>
<th>Morning Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n Control (mmHg)</td>
</tr>
<tr>
<td>Captopril 3</td>
<td>9 5.2 ± 1.3</td>
</tr>
<tr>
<td>Captopril 10</td>
<td>8 4.9 ± 0.7</td>
</tr>
<tr>
<td>Captopril 30</td>
<td>9 4.5 ± 0.5</td>
</tr>
<tr>
<td>Enalaprilat</td>
<td>8 5.5 ± 1.2</td>
</tr>
<tr>
<td>Losartan</td>
<td>8 4.2 ± 1.4</td>
</tr>
<tr>
<td>Lisinopril</td>
<td>6 6.0 ± 0.6</td>
</tr>
</tbody>
</table>

Values are the average difference (over day 3-5) of the 12-h lights-off and 12-h lights-on period. Values are expressed as mean±SEM. *: lisinopril in drinking water; †: P < 0.05 treatment vs control; †: P < 0.05 morning vs evening dose.

### Table 7.4. Average values and between day variability of trough-to-peak ratios.

<table>
<thead>
<tr>
<th>Evening Dose</th>
<th>Morning Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n T/P (%) CV (%)</td>
</tr>
<tr>
<td>Captopril 3</td>
<td>9 27 ± 13</td>
</tr>
<tr>
<td>Captopril 10</td>
<td>8 18 ± 7</td>
</tr>
<tr>
<td>Captopril 30</td>
<td>9 30 ± 3</td>
</tr>
<tr>
<td>Enalaprilat</td>
<td>8 20 ± 13</td>
</tr>
<tr>
<td>Losartan</td>
<td>8 27 ± 7</td>
</tr>
<tr>
<td>Lisinopril</td>
<td>not applicable</td>
</tr>
</tbody>
</table>

The trough-to-peak (T/P) ratio was calculated in 2 ways. First, it was calculated as the average value over day 3-5 of treatment in each individual rat. The variability in this measure is shown as the coefficient of variation (CV). For a comparison T/P ratio's were calculated from the group averages of peak and trough effects over days 3 to 5. Values are expressed as mean±SEM. †: P < 0.05 morning vs evening dose. For the evening dose of lisinopril no T/P peak ratio was calculated since the drug was administered via the drinking water.
Figure 7.6. Comparison of 24-h differences (drug versus saline) in mean arterial pressure (MAP), heart rate (HR) and heart rate product (HRP) during the five day treatment (D1-D5) with enalaprilat (0.3 mg·kg⁻¹), lisinopril (0.1 mg·kg⁻¹), and losartan (10 mg·kg⁻¹) at a morning (○) or evening (■) dose. ○: average values during the 6 h before the start of day 1. R1, R2: values during the two days of recovery (see figure 7.1). Values are means ± SEM. * P < 0.05 morning versus evening dose.

The light period vary too. As shown in Table 7.3, the mean dark-light difference during saline was about 5 mmHg. During captopril treatment this difference increased to 10 mmHg when the infusions were given during the evening period, but fell to 3 mmHg when the infusions were given in the morning period.

The above described effects of captopril 10 mg·kg⁻¹·6 h⁻¹ were dose-dependent, being smaller and greater when the dose of captopril was 3 mg·kg⁻¹·6 h⁻¹ and 30 mg·kg⁻¹·6 h⁻¹, respectively. The data obtained with these doses are summarized in the Tables 7.2-7.4 and Figure 7.5.

After cessation of the last infusion of captopril, MAP returned to near control values during the dark period on day 6. Then, when the lights went on MAP fell again below control on day 6 although no drug was infused. This effect was absent on day 7, the second day of withdrawal.

**Time-related effects of enalaprilate**

Following treatment with enalaprilate, there was a progressive lowering of MAP over the days, reaching stability after 3 days of infusion (Figure 7.6). The average circadian patterns obtained during the evening and morning infusion with enalaprilat are given in Figure 7.7. The average reductions of blood pressure and increments of heart rate over days 3-5 of treatment were not different for the evening and morning regimen, although there was a tendency for blood pressures being lower during the
Figure 7.7. Effects of treatment with enalaprilat (0.3 mg kg⁻¹) on mean arterial pressure (MAP) and heart rate (HR) over a full 24-hour period. SHR were infused either during the morning (open circles) or evening period (closed circles). Values are means over day 3-5 of treatment. The values obtained during the saline infusions were averaged over the morning and evening groups (straight lines). The number of animals in each group is given in Table 7.1. The closed bars denote the lights-off period of the day.

Figure 7.8. Effects of treatment with lisinopril (0.1 mg kg⁻¹) on mean arterial pressure (MAP) and heart rate (HR) over a full 24-hour period. SHR were infused either during the morning (open circles) or lisinopril was given in equivalent doses in the drinking water (closed circles). Values are means over day 3-5 of treatment. The values obtained during the saline infusions (respectively, water drinking) were averaged over the morning and evening groups (straight lines). The number of animals in each group is given in Table 7.1. The closed bars denote the lights-off period of the day.

morning regimen and reflex tachycardia being greater when the evening dose was applied. The circadian amplitude of MAP did not change during treatment with enalaprilat (Table 7.2). Differences between the 12-h dark and 12-h light period were smaller (nearly significantly P<0.1) when enalaprilat was given as a morning than when it was given as an evening dose (table 7.3).

Time-related effects of lisinopril

In one group of SHR, lisinopril was infused for 5 days in the morning period and in the other it was given in the drinking water for 5 days. During both treatment schedules blood pressure reductions reached a plateau on day 3 of the experiment. The average decrease of 24 h blood pressure values during days 3 to 5 was not different between these two groups (Figure 7.6). However, as shown in Figure 8, the greatest protection during the early morning hours in the rat was obtained with the morning infusion rather than when lisinopril was taken via the drinking water. Heart rate was not significantly altered during both treatment regimens (Figure 7.8). With the morning dose of lisinopril, the maximal amplitude increased (Table 7.2), whereas the difference
between the dark and light period became significantly smaller (Table 7.3). These parameters remained unchanged when lisinopril was given in the drinking water.

**Time-related effects of losartan**

As with captopril, enalaprilate, and lisinopril the blood pressure lowering effect of this agent increased during the days of treatment reaching stability at day 3 (Figures 7.6 and 7.9). During both treatment regimens the average daily reduction in MAP was about 15 mmHg. As shown in Figure 7.9 and 7.10, the main antihypertensive effect of losartan was not confined to the infusion period as with captopril and enalaprilate, but lasted much longer. Especially with the morning dose the 24-h blood pressure patterns remained relatively flat and the rise in blood pressure was blunted during the transition from the light to dark period. Table 7.3 indicates that the average difference of MAP between the dark and light period was significantly smaller for the morning than for the evening dose. The circadian amplitude of MAP was not significantly altered from the values obtained during saline (Table 7.2).

Losartan did not influence heart rate significantly. The higher values of HR following the evening dose are the consequence of higher baseline values of HR in this group (Table 7.1). The average reduction in RPP tended to be greater for the morning than for the evening dose (Figure 7.6).
Comparison of averaged effects over all treatment groups

We compared the overall efficacy of treatment during the morning period with treatment during the evening period by averaging the data over each subgroup. Following the analysis of variance, the decrease in RPP was significantly more pronounced when agents were given in the morning period (-8.7±0.9 %) than in the evening period (-4.1±1.2 %, P=0.003, Figures 7.5 and 7.7). Averaged 24-h reductions in MAP and averaged changes in HR were not different between the morning and evening dosing regimens. Through-to-peak ratio’s are compared in Table 7.4. These ratio’s were significantly higher when captopril 10 mg·kg⁻¹ and losartan were given as a morning than evening dose. In general the values were not higher than 50%. The within animal variability in the trough-to-peak ratio, expressed as the coefficient of variation, was in all groups about 100%. To reduce the variability we calculated the trough-to-peak ratio also from the average trough and peak values of blood pressure in each group. In this case, the trough-to-peak ratio’s were about 15% higher. Upon withdrawal of all agents rebound hypertension was never observed.

Discussion

The main finding of this study is that with timed administration of inhibitors of the renin-angiotensin system the circadian blood pressure rhythm of SHR can be modulated to reduce the early morning blood pressure surge. Although, in general, the average reduction over 24 hours was not different between the morning or evening treatment regimen, the blood pressure patterns were different between the two regimens during specific periods of the day. With the morning dose differences between day-night blood pressures were smaller than with the evening dose, whereas with the latter regimen often reflex tachycardia occurred during the sleeping period. The rate pressure product, an index of cardiac oxygen consumption, was lower following the
morning than evening dose. With the short-term acting agents captopril and enalaprilat, blood pressure returned at the end of the daily dosing interval to control levels, whereas this effect was less pronounced following the long-term acting agents lisinopril and losartan. Upon withdrawal of all agents, blood pressure returned gradually to control. Rebound hypertension was never observed.

Run-in effects, pharmacokinetic and pharmacodynamic mechanisms.

One of the remarkable findings of the present study is that the blood pressure lowering effect increased when the ACE inhibitors and losartan were administered repeatedly over 5 days. It may be common knowledge that upon starting antihypertensive treatment so-called run-in effects occur. Especially in the design of clinical studies one accounts for such effects, however, this is the first time that they are shown to occur in rats. In general, in SHR, stable blood pressure reductions were obtained after the third dosing interval. The only exception to this was the repeated morning infusion of 10 mg.kg⁻¹ of captopril. In this case, the antihypertensive effect increased over all days of treatment (Figure 7.3). The latter effect was not drug or dose-dependent since stable effects were reached after 3 days with the 3 and 30 mg.kg⁻¹ dose of captopril. Several mechanism may account for the increasing efficacy over days of treatment. For the agents with a longer half life, such as lisinopril and losartan, it is reasonable to assume that the degree of ACE inhibition and AT₁ receptor blockade, respectively, increases over the days due to accumulation of the drug or its active metabolites. For a short-acting agent like captopril, it seems rather unlikely that this occurs because then the average reduction in blood pressure on the first day of infusion should be much greater for the 30 mg.kg⁻¹ than for the 3 mg.kg⁻¹ dose. The plasma half life of captopril and the terminal half life of the enzyme inhibitor complex are shorter than 2 hours. Consequently the amount of plasma ACE inhibition after 24 hours is probably minimal, which could be the explanation that blood pressure returned to near control values at the end of each dose interval. For both enalaprilat and lisinopril the terminal half life of the enzyme inhibitor complex is much longer (about 20 h). Nonetheless, also for enalaprilat blood pressure returned close to control values at the end of each dose interval. Other factors such as differences in tissue penetration and deposition or differences in paracrine or autocrine actions, may explain this between drug variability.

Irrespective of such variable pharmacokinetics, it is likely that a common additional mechanisms is involved because the dynamic changes on blood pressure were very similar for all ACE inhibitors. From the blood pressure patterns one may deduce that the repeated infusion of ACE-inhibitors has sensitized the cardiovascular system for ACE inhibition. ACE inhibitors have natriuretic properties and the loss of salt may have increased the extent to which blood pressure becomes dependent upon renin or angiotensin II (AII). This dependency is likely to be highest at the end of each dose interval and consequently, the next dose of the antihypertensive agents leads to greater fall in blood pressure. The fact that ACE inhibitors do have greater antihypertensive properties during low salt intake corroborates this hypothesis.

Also with losartan similar run-in effects occurred. At peak concentrations the dose of losartan (10 mg.kg⁻¹) is supposed to block about 90% of all AT₁ receptors and shifts the AII pressor response curve about a factor 35 in rats. With this dose, the degree of AT₁ receptor blockade is probably not very different between day 1 and day
Circadian-phase dependent mechanisms

Mechanisms responsible for differences in the blood pressure patterns between the morning and evening treatment regimen with the ACE inhibitors and losartan must be related to a circadian variation in one or more of the components of the renin-angiotensin system. Both circadian phase-dependent differences in pharmacokinetics as well as pharmacodynamics may be involved. In the present study we have not examined pharmacokinetic data concomitantly with the blood pressure measurements. Because of the limited amount of blood in rats, it is impossible to obtain blood samples repeatedly, without interfering with the blood pressure measurements. Moreover, as discussed below, several findings disprove an important role for circadian phase-dependency of pharmacokinetics.

1. Role of AII

The difference in the blood pressure patterns between the morning and evening dose must be related to time-dependency of the production of AII, since comparable time-dependent patterns were observed with the AT_1 receptor antagonist losartan and ACE inhibitors. Furthermore, this finding excludes an important role for kinins in causing these time dependent effects. This latter aspect is supported by a recent study in TGR in which blood pressure reductions following enalaprilat and losartan were comparable. A specific effect of brain AII can be ruled out too. The time-dependent antihypertensive patterns of captopril were not different from those of enalaprilat, an agent which does not readily cross the blood brain barrier. To which extent AII fluctuates over 24-h periods in non-treated SHR is unknown. In one study in rats, peak values were found in the early morning, coinciding with those of aldosterone. Following the same way of reasoning as above, throughout the 24-h cycle, blood pressure may be most dependent upon AII when sodium intake is lowest, i.e. at the end of the resting period just before resuming feeding and activity. This could be one of the major reasons to explain the efficacy of losartan during this period of the day, i.e. the early morning period (see Figure 7.19).

2. Role of ACE

The circadian variation in the amount of circulating ACE is probably not responsible for the time-dependent efficacy of the present antihypertensive. In SHR, plasma ACE concentrations were relatively constant throughout the 24-h period. Also tissue ACE did not vary markedly over 24 hours. However, in the latter case only data about ACE concentrations in the lungs were studied. Therefore, circadian variations in tissue ACE of organs, which play a major role in systemic resistance to blood flow, cannot be excluded. Also in healthy subjects pharmacokinetics of enalapril were not different after an evening or morning dose.
Chapter 7

3. Role of renin

The most likely rate limiting factor responsible for the time-dependent effects of these drugs is renin. Under normal conditions, in rats, plasma renin levels increase from a trough value at the onset of the resting period to reach maximal values at the time of awakening.\textsuperscript{19, 20} This pattern mirrors the circadian variation in blood pressure, which suggests that, especially during the resting period, renin release may be largely dependent on the renal perfusion pressure.\textsuperscript{24} On the other hand, the circadian rhythm of renin may depend on the sympathetic nervous activity to the kidney\textsuperscript{14} because peak levels of noradrenaline and renin coincide in the early morning hours. This view is supported by the observation that the circadian periodicity in plasma renin levels disappears during chronic \textbeta-blockade in patients as well as after bilateral nephrectomy.\textsuperscript{21} Furthermore, kidney-derived renin is known to be rate-limiting in the production of AII by tissues.\textsuperscript{15} According to this, the antihypertensive effect of the early morning infusion of the ACE inhibitors may have been caused by blunting the peak values of renin either via renal vasodilation or sympathetic inhibition. Clearly, the latter mechanism is involved because following the morning doses of the ACE inhibitors, the degree of reflex tachycardia was less than following the evening dose.

An implication of renin being possibly the rate-limiting factor is that the effect of ACE inhibition on blood pressure is low or absent when the production of renin is low.\textsuperscript{19} In rats, this is most-likely to occur at the end of the active period, when they ingest a significant amount of food to bridge the resting period. At this time point plasma renin is lowest\textsuperscript{19} and we found least antihypertensive efficacy. Also the blood pressure pattern obtained during the 2 days of recovery support that the circadian variation in one of the endogenous components is rate limiting for the action of the ACE inhibitors and losartan. Upon withdrawal of the last morning dose of captopril, blood pressure returned to control values. However, then during the progression of the resting period, blood pressure fell again below control levels. A similar secondary fall in blood pressure in the resting period effect was observed after the last morning dose of losartan (Figure 7.10). This pattern was absent after the last evening dose. Similar delayed effects have been reported to occur in patients too. Smith et al.\textsuperscript{22} showed that after withdrawal of candesartan when blood pressure had returned to control values between 14.00 and 24.00 h, it fell again below control in the ensuing night between 0 and 08 h. Based on these data we suggest that the physiological production of renin is determining the circadian variation in the antihypertensive efficacy of the ACE inhibitors and losartan.

Trough-to-peak ratio

In the current study we have examined the effect of the dosing time on the trough-to-peak ratio of the ACE inhibitors and losartan. For captopril (10 mg.kg\textsuperscript{-1}) and losartan the trough-to-peak ratio was significantly higher when the drugs were infused during the morning. However, only for the morning dose of losartan the trough-to-peak ratio was above 50 % and in the range (50%-66%) required for FDA approval.\textsuperscript{11, 12} The other drugs failed to meet this criterion. Obviously species differences may explain why these values are lower in rats than in humans. Because of a high metabolic rate in rats, the half-lives of the present agents are probably shorter than in humans. However, it is of our concern that even in our controlled experimental setup with continuous
monitoring of blood pressure, the within animal variability of the trough-to-peak values was very high (about 100%). A clear explanation for the trough-to-peak values being so inconstant is lacking. Factors involved are probably the day-to-day variations in blood pressure associated with the activity/sleep pattern and the day-to-day variation in renin release which occurs burst like especially during sleep[1]. Our present data in rats add to the concerns formulated by others on the employment of this index[12,37].

Therapeutic implications

In the current study we have shown that blood pressure patterns in hypertensive animals can be modulated with timed dosing of drugs that antagonize the renin angiotensin system. So far 4 studies in patients have been conducted[7-10]. Although they disagree to a certain extent about the optimal time period of administration, all these studies showed that the timing of the administration of ACE inhibitors was important for the ensuing circadian blood pressure pattern. Which 24-h blood pressure pattern has an optimal protection for the morbid events related to hypertension has to be established in clinical trials. Our data in SHR, foster the development of ACE preparations or AT1-receptor antagonist which also have a controlled onset to anticipate upon the circadian variations in components of the renin-angiotensin system.

In conclusion, circadian blood pressure rhythms in SHR were differentially modulated by a circadian-phase dependent administration of ACE inhibitors and the AT1-receptor antagonist losartan. The differential effects were more likely due to circadian variations in components of the renin angiotensin system than circadian differences in the pharmacokinetics of these agents.

References

Chapter 7


Chapter 8
General discussion
In this thesis we have investigated cardiovascular regulatory mechanisms in hypertension, and the effect of anti-hypertensive treatment in relation to the time of day. For this reason we focussed in detail on the short-term and long-term variability of time-series of cardiovascular parameters. The aim of this final chapter is to place the present findings in perspective, emphasizing on the pathophysiological consequences of the circadian patterns of hemodynamics in hypertension. Finally, the implications for therapy are discussed.

**Time-series analysis**

Rhythmic variability can be used as a non-destructive probe of regulatory mechanisms. Spectral analysis of heart rate and, to a lesser extent, of blood pressure have been used to quantify autonomic function in a number of pathophysiological states. Fast fluctuations in heart rate related to respiration are mainly associated with parasympathetic activity, whereas slower oscillations (the so-called Mayer waves around 0.1 Hz) are mostly attributed to sympathetic activity.

In order to make judgements on the activity of a regulatory mechanism based on spectral analysis one should be sure that there is only one mechanism that has the ability to cause oscillations in that particular frequency range. In chapter 2 we have measured beat-to-beat variability of regional blood flows and cardiac output to determine the hemodynamic origin of oscillations in blood pressure. Spectral analysis showed that high frequency, respiratory related, fluctuations of blood pressure were mechanically coupled to variability of cardiac output, i.e. mainly stroke volume. Blood pressure oscillations in the lower frequency ranges resulted mainly from myogenic-like oscillations in regional blood flows and not from autonomic nervous activity. In addition, it has recently been demonstrated that also endothelial derived factors contribute to fluctuations of blood pressure in the low frequency range. Thus, in the low frequency area at 0.1 Hz blood pressure variability is influenced by many factors. In contrast, heart rate fluctuations in this frequency range seem to follow the blood pressure variations and may mainly reflect baroreflex activity. In chapter 3 we have used spectral analysis in our search for a on-line method to derive baroreflex sensitivity from beat-to-beat changes in blood pressure and heart rate. In our rat strains, however, the values obtained from these calculations showed only little correlation with baroreflex gains measured with the classic pharmacological method. For testing the autonomic state in patients, classic tests like deep breathing and Valsalva’s manoeuvre usually suffice. When autonomic function must be monitored for longer periods, spectral analysis of heart rate signals and blood pressure can be considered. However, these data should be interpreted with care. Up to now, there are no data bases with values of spectral measures obtained in a normal healthy population. On the other hand, there are striking similarities in spectral measures of blood pressure between SHR and essential hypertensive patients over 24-h periods. In both species, low frequency oscillations in blood pressure are increased and high and low oscillations of heart rate are decreased. Also baroreflex gains over 24 hour periods as calculated by sequential methods revealed similar results in SHR and essential hypertensive patients. Thus, whereas these tools seem to have their diagnostic value, the underlying pathophysiological mechanisms are not fully clear.

The present thesis stresses the importance of measuring cardiovascular function
throughout the day. In chapter 4 we found an intriguing inverse effect between the circadian and ultradian effects of total autonomic blockade on variability of blood pressure and heart rate. Total autonomic blockade produced the expected decrease of short-term variability of heart rate and a slight increase in short-term variability of blood pressure. The opposite was true for the circadian variability of these parameters. The 24 h heart rate rhythm remained unaltered, whereas the 24-h rhythm of blood pressure was abolished. A comparable phenomenon was seen in humans after cardiac transplantation. Thus long-term variations in heart rate seem to be rather independent of autonomic tone and may be more related to intrinsic mechanisms such as atrial filling.

Hemodynamic monitoring over prolonged periods of time poses the question how the data must be gathered. Rats have almost half a million heart beats every day, and data reduction is necessary to perform statistical analyses on the results. However, this process carries the risk of losing important information or interesting phenomena hidden in short-term sequences of these parameters. Modern computers are fast enough to compute on-line measures of the dynamics of the cardiovascular system (see chapter 3). We believe, that the application of such techniques will lead to new diagnostic tools and add to our understanding and of the cardiovascular system in many pathologies.

In addition to these time-related methods we have used 2-dimensional frequency distributions of beat-to-beat values of complete 24 hour periods. This was done to gain an insight into the distribution of hemodynamic parameters, together with the beat-to-beat interactions between 2 parameters. Using this technique we could show (chapter 5) that on a circadian time-scale variations of cardiac output were greater than variations of blood pressure. This suggests that even on such a long-term time-scale blood pressure is controlled tighter than cardiac output. In our view there is no such thing as a set-point for cardiac output. The adaptive processes that control cardiac output over extended periods, such as Guyton's whole-body autoregulation, occur in our view at organ level. Over long-term periods, every vascular bed regulates its own flow as a function of its requirements. Because these requirement vary throughout the day, relatively large variations are observed. The application of the 2-dimensional frequency distribution also showed that, unlike values of other systemic hemodynamic parameters, heart rate values were clustered at certain preferred levels. It seems that, at least in rats, the heart likes to work in certain gears. The underlying mechanism of this phenomenon is unknown. Certain heart rates could be energetically more effective, or preferred by internal properties of the pacemaker cells in the sinus node. In conclusion, the calculation of measures derived from beat-to-beat values gives us an unique insight into the dynamics of cardiovascular function over prolonged periods of time, although more work is needed to validate such probes clinically.

**Circadian cardiovascular function and pathophysiology**

Epidemiological studies have shown that the onset of several cardiovascular morbid events occurs more often in the early morning. The circadian pattern of several physiological functions may explain the striking of such events at this time-point of the day. First it must be stated that the vascular system in these patients is already vulnerable, mostly by the occurrence of atherosclerosis in coronary arteries or arteries providing the brain with blood. The physical stress that eventually causes the injury to the vessel wall is not evenly distributed throughout the day. It has already been well
established that blood pressure and heart rate are higher during the active period of the day. The increase in blood pressure in the morning causes an increase in the passive strain on the arterial system, while the increase in heart rate causes an additional increase of the dynamic strain. The concomitant increase of rate pressure product augments the workload of the heart, and increases oxygen consumption. Cardiac output is also higher during the active period, but the data on the circadian pattern of total peripheral resistance are inconsistent. This may be due to methodological problems. In chapter 5 we have made direct measurements of cardiac output with electromagnetic flowprobes in unanesthetized rats over 24 hours. In that study we have shown that total peripheral resistance is higher during the passive phase of the day, because the relative circadian variation in cardiac output is higher than the relative circadian variation in blood pressure. Until recently it has been difficult to measure cardiac output in humans without stressing the subject under study. The pulse contour and Modelflow method have been developed to measure an indirect index of cardiac output from peripheral blood pressure recordings in humans. These methods assure the undisturbed continuous measurement of cardiac output. Through these methods it has been assessed that also in humans total peripheral resistance is higher in the passive period. We have also shown that total peripheral resistance is still high at the transition from the sleeping to the active stage, and only starts to decrease when the animals become active. This imposes an increased afterload in the morning. The heart has to pump harder in order to ensure sufficient flow against a high resistance. An extra factor in keeping resistance high in the morning could be the impairment of flow-induced vasodilatation in hypertension. In chapter 4 we have shown that baroreflex function is decreased at this time point, causing an increase in short-term blood pressure variability. In short, heart rate, arterial pressure, blood pressure variability, and peripheral resistance all contribute to the high stress on the arterial wall during the early morning. The increase in stress on the wall could lead to damage of predisposed or fragile sites like atherosclerotic plaques. In particular, the rupture of these plaques is now recognized as a major pathogenic mechanism for myocardial or brain infarcts. Other factors like an increased susceptibility for the generation of blood clots at this time-point even increase the chance that an atherosclerotic rupture leads to a life threatening infarction.

The incidence of myocardial infarction also shows a weekly pattern, with the highest incidence on Mondays. This phenomenon can be explained by the tendency of humans to sleep out during the weekend. This causes a phase-delay in the endogenous regulatory systems controlling blood pressure. When the activity pattern of weekdays is resumed on Monday morning, a phase-advance is required, whereas the regulatory systems are still geared for inactivity, with high peripheral resistance. Blood pressure and heart rate will have to increase more than normally to guarantee sufficient blood flow. This can be viewed as an exaggerated form of the normal adaptations to the circulation in the early morning.

Circadian considerations for therapy

Much has been done in the last 40 years to improve anti-hypertensive therapy. From the early sympatholytic agents to modern drugs like ACE-inhibitors and calcium antagonists, anti-hypertensive agents have become safer and more devoid of adverse
effects. Yet anti-hypertensive treatment is not yet optimal. People treated for hypertension still have a higher risk for diseases associated with hypertension than people with normal blood pressure. 27-30. A number of factors can be held responsible for this failure of anti-hypertensive treatment [see Introduction]. One of these factors is the observation that current anti-hypertensive therapy does not take the circadian rhythmicity of cardiovascular events into account.

To understand the time-related effects of anti-hypertensive treatment it is important to know the major determinants of circadian patterns of hemodynamics. Physical activity is the most important factor for the level of blood pressure and heart rate, but it is likely that there is also an endogenous blood pressure rhythm. 29. Of the regulatory mechanisms it seems that the autonomic nervous system has a causal role in the generation of the circadian pattern of blood pressure. 32. The balance between parasympathetic and sympathetic nervous system is shifted towards the sympathetic side during the active period and towards the parasympathetic side during the passive period. 33-36. In contrast to the autonomic nervous system, other regulatory mechanisms, like the renin angiotensin system and vasopressin show adaptive or counter regulatory behaviour on the circadian timescale, with higher activity during the passive period. 35, 37-40. This phenomenon is further illustrated by the reversal of the circadian pattern of blood pressure in certain (renin-dependent) secondary forms of hypertension and in Renin-2 transgenic rats. 41. The circadian patterns of these hormones can explain the increase in total peripheral resistance during the passive period. These differences in regulatory mechanisms, may imply that drugs targeted at the sympathetic nervous system, like &alpha;1- and &beta;-blockers blunt the circadian variability of blood pressure, whereas drugs targeted at the renin angiotensin system can increase circadian variability of blood pressure, and may be used to restore the decrease of the circadian blood pressure variability in non-dippers.

Current anti-hypertensive treatment is primarily aimed at providing an even blood pressure reduction throughout the day. 42. Many studies have examined the 24-hour efficacy of anti-hypertensive agents (for reviews see). 43-45. A general finding of these studies is that, although anti-hypertensive treatment reduces the average blood pressure level, the circadian pattern of blood pressure is preserved and that treatment does not attenuate the early morning rise in blood pressure. 44. It is yet unclear what the goal should be in a circadian approach towards anti-hypertensive treatment. Sirgo et al. 46 have stressed the importance of anti-hypertensive efficacy of drugs all through the day with an emphasis on the early morning, which is often situated at the end of a dosing interval. This evenly distributed effect throughout the day is also part of the recommendations of the FDA for new drugs, and is generally achieved with long-acting agents. In hypertensive subjects where the normal difference between the active and the resting period is diminished, or even reversed, an increased incidence of end organ damage is seen, 46, 47, and anti-hypertensive treatment fails to restore the normal difference between nighttime and daytime blood pressure. Here the timing of dosing could be used to restore the normal day-night pattern of blood pressure.

The FDA has published guidelines for the registration of new anti-hypertensive drugs. 48. One of the most controversial of these guidelines was the requirement to have a trough to peak ratio of at least 50% when using the recommended dosing scheme. 49. The trough to peak ratio was designed to ensure efficacy at the end of the dosing
interval, without having to resort to inappropriately high doses. In the chapter 7 we have examined the effect of the dosing time on trough to peak ratio. In general the trough to peak ratio of blood pressure was higher when the drugs were infused during the morning. Only in one case (losartan morning dose) the trough to peak ratio was above 50%. The other drugs failed to meet the cut-off point. The high metabolic rate of rats, and therefore shorter half-life of drugs, could play a role here, as well as the fact that blood pressure was recorded continuously in contrast to most clinical studies where blood pressure is measured in intervals. However, it is of our concern that even in our highly controlled experimental setup with continuous monitoring of blood pressure, we were not confident of the obtained values. The inter individual variability of trough to peak ratio was unacceptably high, with a coefficient of variation of about 100% in most groups, and large differences were found between two ways to calculate trough to peak ratio. These factors should lead to reconsidering the value of this parameter.

We believe that restoring the circadian blood pressure pattern to normal may not be sufficient in this group of patients with an increased risk for cardiovascular illness. We hypothesize that it could be beneficial to desynchronize the circadian patterns of the processes that now all show their peak stress in the early morning. One way of doing this is by delaying the rise in blood pressure at this moment by treating the patient with short-acting drugs in the early morning. In chapter 7 we have shown that it is possible to manipulate blood pressure patterns in this way. In humans the exact timing at this period is difficult because the patients are still asleep; at the moment the drugs should have their peak effect. However, release preparations have been devised in the meantime, that are specifically targeted at this period. The approach of changing the pattern of blood pressure will require treatment with the use of short-acting drugs. This is in contrast with the current tendency to prolong the half life of anti-hypertensive drugs, thus obtaining a flat anti-hypertensive profile.

A serious drawback of therapy aimed at changing the pattern of blood pressure is that it requires the patient to take medication at the same time every day. Compliance is low for anti-hypertensive medication, whereas strict adherence to the dosing schedule is important for chronopharmacology. Also in order to monitor the treatment it will be necessary to carry out regular 24 hour ambulatory blood pressure measurements. This can lead to therapy based on risk-stratification. People at a higher risk for adverse cardiovascular events, with other risk factors like old age, end organ damage, left ventricular hypertrophy, obesity, and/or smoking, will require more aggressive therapy of hypertension, whereas low-risk patients probably will be sufficiently treated with current anti-hypertensive therapeutic strategies.

Currently most attention is given to blood pressure levels, whereas other hemodynamic parameters are more or less neglected. Our study in chapter 7 showed that when the dose was targeted at different time points of the day, not so much the blood pressure reduction was different between time points, but the heart rate. This can be an important consideration for treatment, because the workload of the heart is rate dependent. At the moment there is insufficient data to interpret circadian patterns of peripheral resistance and cardiac output during treatment, although it is conceivable that especially drugs targeted at the renin angiotensin system can be beneficial in lowering total peripheral resistance in the morning.
Concluding remarks

Despite the reduction of blood pressure the incidence of cardiac life threatening events is still higher in treated hypertensive subjects than one would expect on the basis of the achieved reduction of blood pressure^20.30.

We have shown that with a chronotherapeutic approach to anti-hypertensive treatment it is possible to change the pattern of blood pressure throughout the day, thereby desynchronizing the circadian patterns that exert a physical stress on the arterial wall. Prospective trials in humans will be necessary to elucidate if a chronotherapeutic approach to hypertension is beneficial in preventing cardiovascular disease, compared to current anti-hypertensive treatment.

References


Chapter 8.
Chapter 9
Nederlandse samenvatting
Nederlandse samenstelling

Het blijkt dat cardiovasculaire incidenten zoals myocardinfarct, herseninfarct of plotselinge dood vooral voorkomen in de vroege ochtend. Het doel van dit proefschrift is een studie naar de mechanismen die aan dit fenomeen ten grondslag liggen.

Hypertensie is de belangrijkste risicofactor voor het ontstaan van atherosclerose, en daarmee voor het ontstaan van hartinfarcten. In 90% van de patiënten met hypertensie wordt bij onderzoek geen oorzaak voor het ontstaan van hypertensie gevonden. Deze vorm heet primaire of essentiële hypertensie. In de resterende 10% worden afwijkingen van nieren, bijnieren of de nierarterieën gezien. Deze vormen worden secundaire hypertensie genoemd. Het blijkt dat bij primaire hypertensie het dag-nacht ritme van de bloeddruk in het algemeen aanwezig is, terwijl bij secundaire vormen van hypertensie vaak een klein verschil of zelfs een negatief verschil tussen dag- en nachtwerten van de bloeddruk wordt gezien. Patiënten met een klein verschil in bloeddruk tussen dag en nacht (non-dippers) hebben een vergrootte kans op complicaties van hypertensie, zoals linkerventrikel hypertrofie of retinopathie.

De huidige behandelingen richtlijnen voor hypertensie zijn erop gericht de diastolische bloeddruk onder de 95 mmHg te krijgen. De laatste jaren gaan er echter ook stemmen om op patiënten te behandelen afhankelijk van de aanwezigheid van andere risicofactoren als cholesterol plasma spiegels, lichaamswaardig en al aanwezige schade tengevolge van hypertensie. Gezien de hoge incidentie tijdens de ochtend van het voorkomen van myocardinfarct zou het wenselijk zijn dat anti-hypertensieve therapie ook gericht is op de vroege ochtend.

Het onderzoek in dit proefschrift beschrijft in de hoofdstukken 1 tot 5 een aantal fysiologische processen die op de korte en lange termijn een rol spelen bij de regulatie van het cardiovasculaire systeem. In de hoofdstukken 6 en 7 wordt de invloed van een aantal antihypertensieve farmaca op het dag-nacht ritme van cardiovasculaire parameters onderzocht.

In de algemene introductie wordt de fysiologie beschreven van het systeem dat dag-nacht verschillen kan veroorzaken van allerlei processen in het lichaam. Hierbij worden 2 verschillende vormen onderscheiden: 1) Endogene ritme: processen die afhankelijk zijn van een innerlijke klok. Deze processen verloren ook een dag-nacht ritme als er geen dag-nacht verschillen meer zijn in de omgeving. 2) Exogene ritme: processen die dag-nacht verschillen vertonen als reactie op dag-nacht verschillen in de omgeving. Het voordeel van een endogene ritme is dat een proces zich kan instellen op veranderingen die nog aan komen. De meeste fysiologische processen en parameters vertonen dag-nacht verschillen, maar in het grootste deel gedaan is niet bekend of dit door een endogene of exogene ritme bepaald wordt.

Ook binnen het cardiovasculaire systeem volgen de meeste parameters een dag-nacht ritme. Van de bloeddruk en de hartfrequentie is bekend dat zij hogere waarden vertonen gedurende de dag dan opzicht van de nacht. Hartminutvolume en totale perifere weerstand zijn minder goed onderzocht. Voor het hartminutvolume zijn er aanwijzingen dat het overdag hogere waarden vertoont dan 's nachts. De relatie tussen verschillen in hartminutvolume zijn groter dan die van de bloeddruk. Daarom is 's nachts de totale perifere weerstand hoger dan overdag. De invloed van cardiovasculaire regelmechanismen vertoont ook een dag-nacht ritme, waarbij opvatting dat behalve (nep)adrenaline de meeste hormonen met een vasoconstrictieve werking een ritme hebben met hogere plasma spiegels tijdens de nacht.
Er is een aantal fysiologische processen die aanleiding kunnen geven tot een verhoogd risico op cardiovasculaire incidenten tijdens de vroege ochtend. De bloeddruk en de hartfrequentie stijgen gedurende deze periode. Samen met een verhoogde bloeddruk variabiliteit zorgt dit voor een verhoogd fysiologische belasting van de vaatwanden het hart. Bloed stolt makkelijker tijdens deze periode terwijl bloedstolsels minder snel worden afgebroken. Tenslotte toont het hormoon cortisol in deze periode zijn hoogste waarden waardoor coronaire arterieën extra gevoelig zijn voor catecholamines.

In hoofdstuk 2 zijn de korte termijn schommelingen van de bloeddruk vergeleken met schommelingen in het hartminuutvolume en de doorstrooming van nieren, darmen en het achterlijf van niet-geneestheeseerde rustende ratten. Hierbij zijn drie frequentiegebieden onderzocht. 1) Hoog frequent (HF; rond 1.6 Hz): schommelingen rond deze frequentie worden meestal in verband gebracht met de invloed van het parasympathische autonome zenuwstelsel en de ademhaling. In het HF gebied wordt gezien dat de variabiliteit van het slagvolume de variabiliteit van het hartminuutvolume bepaalt, en dat er een directe, mechanische, koppeling is tussen het hartminuutvolume signaal en het bloeddruk signaal. De HF schommelingen worden gedempt voortgeleid naar de perifere vaatbedden van nieren, darmen en achterlijf. 2) Midden frequent (MF; rond 0.4 Hz): deze schommelingen zijn meestal afhankelijk van het functioneren van het autonome zenuwstelsel. MF schommelingen in de bloeddruk worden vooraf veroorzaakt door doorstromingsschommelingen in het achterlijf terwijl de doorstrooming van de nieren en de darmen de schommelingen in de bloeddruk passief volgen. 3) Laag frequent (LF; rond 0.12 Hz): de oorsprong van deze schommelingen dient gezocht te worden in myogene fluctuaties van arterieën in lokale vaatbedden. De LF schommelingen van bloeddruk lijken te worden veroorzaakt door myogene reacties in de vaatbedden van de nieren en de darmen. Het feit dat de myogene reactie een ritme veroorzaakt in de bloeddruk toont aan dat de spieractiviteit van de arterieën tijdens deze reactie gecoördineerd wordt.

In hoofdstuk 3 is een methode ontwikkeld om de baroreceptor reflex gevoeligheid (BRS) bij ratten te meten met behulp van spontane fluctuaties van bloeddruk en hartperiode. Bij vergelijking van de nieuwe methode met een standaard methode bij verschillende omstandigheden blijkt dat de nieuwe methode een goede overeenkomst geeft met de standaard methode.

Het grote voordeel van de in hoofdstuk 3 ontwikkelde methode is dat de BRS continu in vrij bewegende dieren kan worden bepaald. Deze methode is in hoofdstuk 4 gebruikt om te bepalen of de BRS een dag-nacht ritme vertoont. Daarnaast zijn schommelingen in BRS, bloeddruk en hartfrequentie onderzocht met een periode tussen 5 minuten en uren (afwisselende ritmen). Deze onderzoeken zijn gedaan bij normotensieve en hypertensieve ratten. Tevens is de invloed van verschillende middelen onderzocht die het autonome zenuwstelsel beïnvloeden. Met betrekking tot de BRS werd gevonden dat deze gedurende de gehele dag lager is in hypertensieve ratten, vergeleken met normotensieve ratten. Opvallend is dat de BRS zijn minimaal waarde bereikt tijdens de overgang van de slaap naar de actieve periode, de periode waarin bij mensen het voorkomen van myocardinfarcten is verhoogd. Het dag-nacht ritme van de bloeddruk is sterk afhankelijk van de beïnvloeding van het autonome zenuwstelsel. Na uitschakeling van de fysiologische autonome zenuwstelsel op farmacologische wijze is er geen dag-nacht verschil meer in de bloeddruk. Remming van het sympathisch autonoom zenuwstelsel
van de bloedvaten leidt zelfs tot een lagere waarde van de bloeddruk tijdens de actieve periode. Spectrum analyse laat een zogenaamd 1/f verband zien tussen spectrale vermogen en de frequentie. In het algemeen duidt een dergelijk 1/f verband op het ontbreken van een dominant regelmechanisme en de aanwezigheid van meerdere regelmechanismen met invloed in het onderzochte frequentie gebied. Manipulatie van het autonome zenuwstelsel heeft relatief weinig effect op deze relatie. De hartfrequentie reageert zowel voor wat betreft het dag-nacht ritme als de ultradiepe ritmes anders dan de bloeddruk. Het dag-nacht ritme van de hartfrequentie is weinig gevoelig voor verandering van het autonome zenuwstelsel, en lijkt daarmee voornamelijk afhankelijk van intrinsieke cardiale mechanismen.

In hoofdstuk 5 is het dag-nacht ritme van het hartminuutvolume onderzocht, en de invloed die het autonoom zenuwstelsel en de aanwezigheid van hypertensie erop heeft. Ook zijn ritmen met een kortere periode en de slag-op-slag relatie tussen verschillende cardiovasculaire parameters onderzocht. Het blijkt dat het hartminuutvolume tijdens de actieve periode een hogere waarde heeft dan tijdens de slaap periode, en dat de waarden en het dag-nacht patroon bij de normotensieve en hypertensieve ratten vrijwel identiek zijn. Het grote verschil tussen normotensieve en hypertensieve ratten is de totale perifere weerstand die bij hypertensie verhoogd is. Het patroon is echter vergelijkbaar. Na blokkade van het cardiale autonome zenuwstelsel is het hartminuutvolume gedaald, en is het verschil tussen dag en nacht sterk verminderd. Het relatieve vermogen van het hartminuutvolume is ongeveer 2 keer zo groot als die van de bloeddruk in het onderzochte frequentiebereik. Net als de bloeddruk en hartfrequentie in hoofdstuk 4, laten het hartminuutvolume en de totale perifere weerstand een 1/f relatie zien tussen vermogen en frequentie in het frequentiebereik tussen enkele minuten en uren. Er zijn geen verschillen tussen normotensieve en hypertensieve ratten. De steilheid van de 1/f relatie van de weerstand is verhoogd na autonome blokkade, waarschijnlijk als gevolg van het ontbreken van de bufferende werking van de baroreflex. Uit de directe slag-op-slag vergelijking over 24 uur tussen bloeddruk en hartminuutvolume blijkt dat over deze periode het hartminuutvolume meer variabel is dan de bloeddruk, en ook dat er geen rechtsstreeks verband is tussen deze twee parameters. Tussen de perifere weerstand en hartminuutvolume bestaat wel een rechtsstreeks relatie. Omdat de variabiliteit van hartminuutvolume in het hele bereik van hoogfrequentie schommelingen tot het dag-nacht ritme hoger is als die van de bloeddruk lijkt het waarschijnlijk dat dit niet een centraal geregeld parameter is. De figuren van 24-uuurs slag-op-slag relaties tussen hartfrequentie en andere parameters laten regelmatig meerdere pieken zien. Aangezien deze pieken in aantal toenemen na cardiale blokkade is het waarschijnlijk dat deze door een intrinsieke mechanisme van het hart veroorzaakt worden.

In hoofdstuk 6 is het effect van verschillende tijden van toediening bekeken van een aantal klassen van anti-hypertensieve stoffen. Er zijn duidelijke verschillen in effectiviteit en werkingsduur gevonden. De sympathicotyse stoffen zijn het meest effectief in de periode dat de bloeddruk stijgt terwijl renale vaatverwijzers als de ACE-remmers captoprill en de beta-blokkeerder tertatolol meer effectief zijn bij toediening tijdens het begin van de slaap.

In hoofdstuk 7 is een aantal stoffen die het renine-angiotensine systeem beïnvloeden met elkaar vergeleken. De stoffen zijn in deze studie op 5 achtereenvolgende
dagen als infuus gegeven op twee verschillende tijdstippen. De gemiddelde bloeddrukverlaging gedurende 24 uur is niet verschillend tussen een ochtend of een avond dosering, maar er zijn wel duidelijke verschillen in het patroon van de bloeddruk gedurende 24 uur. Het rate-pressure product, een maat voor de zuurstofconsumptie van het hart, daalt tijdens de ochtend dosering sterker dan tijdens de avond dosering. Dit lijkt vooral bepaald door een sterker hartfrequentie verhoging bij de avond dosering. Een interessant fenomeen tijdens deze studie is dat het anti-hypertensieve effect van deze middelen in de loop van een aantal dagen toenaam, zonder dat dit te wijten kon zijn aan ophoping van de stoffen in het lichaam. Het lijkt erop dat de bloeddruk na enkele dagen van behandeling afhankelijk wordt van het angiotensine II, het hormoon dat onderdrukt wordt door deze middelen. De veranderingen in het patroon van de bloeddruk en hartfrequentie geven aan dat de tijdsafhankelijke toediening van ACE-remmers een bijdrage kunnen leveren aan de behandeling van hypertensie.

In dit proefschrift hebben we laten zien dat het mogelijk is om het patroon van de bloeddruk zodanig te veranderen dat de piekbelasting van risicofactoren die belangrijk zijn voor het ontstaan van myocardinfarct niet meer alle op hetzelfde tijdstip van de dag plaatsvinden. In het slothoofdstuk wordt de betekenis van deze observaties besproken in het kader van risicofactoren en behandelstrategieën bij hypertensieve patiënten.
Nederlandse samenvatting
Dankwoord

Het maken van een proefschrift doe je niet alleen. Daarom wil ik op deze plek graag een aantal mensen bedanken die een bijdrage hebben geleverd aan de totstandkoming ervan.

Allereerst Ben Janssen. Beste Ben, zonder jouw was dit proefschrift niet afgekomen. Jouw ideeën zijn de uitgangspunten geweest voor de studies, en dit proefschrift laat ook duidelijk zien dat het onderzoek aan vrij bewegende, niet verdoofde dieren essentieel is in het onderzoek van de circulatie. Je bent steeds in staat geweest om de twijfels weg te nemen die ik had bij het schrijven van de verschillende hoofdstukken. Ik hoop dan ook dat je net zo blij bent als ik, dat het project na al deze jaren toch dit resultaat heeft opgeleverd.


Caroline Eerdmans. Beste Caroline, bedankt voor je hulp op het operatie lab. Je hebt er altijd voor gezorgd dat de dieren voor mijn experimenten perfect voorbereid waren.

Verder wil ik de rest van de vakgroep farmacologie bedanken voor de prettige samenwerking en de goede sfeer.

De medewerkers van de proefdiervoorzieningen, bedankt voor alle inspanningen voor het leveren van de ratten, en de hulp tijdens de verhuizing naar de nieuwbouw, waardoor het mogelijk was mijn experimenten te vervolgen in de kelder.

De instrumentele dienst. Bedankt voor alle snelle hulp bij problemen met het meet programma en het ontwikkelen van de nieuwe meetopstelling.

Ook wil ik graag iedereen bedanken die mijn verblijf in Maastricht zo aangenaam hebben gemaakt. De leden van de schermvereniging MAS in Contro, die doordrongen zijn van de prioriteit die een evaluatie van een training moet hebben. De zwemclub, die Elsbeth en mij bij elkaar gebracht heeft. De inwoners van #descent en #w-e, waar ik mijn eerste schreden op IRC deed.


Als laatste mijn ouders. Pap, mam, ik wil jullie bedanken voor de wijze waarop jullie me altijd hebben gesteund in mijn keuzes.
Curriculum Vitae

1975-1981  Atheneum te Groningen
1981-1989  Studie Geneeskunde te Groningen
1989-1991  Dienstplicht, vervuld in het militair revalidatie centrum te Doorn
1991      Tijdelijk werk in verpleeghuizen te Almen, Deventer en Lelystad
1991-1996  Assistent in Opleiding bij afdeling farmacologie Universiteit
           Maastricht. Project 'Circadiane variabiliteit van bloeddruk regulatie
           mechanismen bij hypertensie'
1997-heden  Arts-onderzoeker bij afdeling Pathologie LUMC Leiden. Project 'Blob
           informatie en Presentatie Systeem.

Publicaties

Full papers

- Janssen BJA, Oosting J, Tyssen CM, Struijk-Boudier HAJ: Time-dependent
  efficacy of antihypertensive agents in spontaneously hypertensive rats. Chronobiol
- Janssen BJA, Oosting J, Slaaf DW, Pesson PB, Struijk-Boudier HAJ:
  Hemodynamic basis of oscillations in systemic arterial pressure in conscious rats. Am
- Oosting J, Struijk-Boudier HAJ, Janssen BJA: Validation of a continuous barorecep-
  tor reflex sensitivity index calculated from spontaneous fluctuations of blood pressure
- Oosting J, Struijk-Boudier HAJ, Janssen BJA: Autonomic control of ultradian and
  circadian rhythms of blood pressure, heart rate, and baroreflex sensitivity in spontane-
- Oosting J, Struijk-Boudier HAJ, Janssen BJA: Circadian and ultradian control of
  cardiac output in spontaneous hypertension in rats. Am J Physiol 1997; 273: H66-
  H75.

Abstracts

- Oosting J, Tyssen CM, Struijk-Boudier HAJ, Janssen BJA; Effects of angiotensin
  converting enzyme inhibitors on the early morning rise in blood pressure in rats; Pharm
  Wld Sci Ed 1992; 14: H10
- Oosting J, Eerdmons-Tyssen CM, Janssen BJA: Baroreflex sensitivity in rats;
  validation of methods using spontaneous blood pressure changes. Eur J Physiol (Pfliegers
  Arch) 1993; 424: R6.
- Oosting J, Struijk-Boudier HAJ, Janssen BJA; Role of the baroreflex in the hypo-
  tensive response following short-term phenylephrine-induced pressor effects. J
- Oosting J, Eerdmons-Tyssen CM, Janssen BJA; Chronopharmacology of Preparanol
  and Captopril in Spontaneously hypertensive rats; Phar Tor 1993; 72: Suppl II # 53
- Oosting J, Janssen BJA; Diurnal pattern of baroreceptor reflex sensitivity in WKY
  and SHR rats; Phar Wld Ser C 1993: 156 Suppl 1
- Janssen BJA, Oosting J, Struijk-Boudier HAJ; Chronopharmacology of
  antihypertensives in spontaneously hypertensive rats; Can J Phys Phar 1994;
  72:Suppl 1;137
- Oosting J, Janssen BJA, Struijker-Boudier HAJ; 24 hour baroreceptor reflex control in WKY and SHR rats; *Canj Phys Phar* 1994: 72:Suppl 1;608
- Oosting J, Struijker-Boudier HAJ, Janssen BJA; Intermittent infusions of vasoconstrictive agents cause hypotension in the drug-free period in SHR; *Phar Wrld Scnc* 1994 16:6 Suppl J;12