Role of the medulla oblongata in normal and high arterial blood pressure regulation: the contribution of Escola Paulista de Medicina – UNIFESP

SERGIO L. CRAVO1, RUY R. CAMPOS1, EDUARDO COLOMBARI1, MÔNICA A. SATO2, CÂSIA M. BERGAMASCHI1, GUSTAVO R. PEDRINO1, MARCOS L. FERREIRA-NETO4 and OSWALDO U. LOPES1

1Departamento de Fisiologia, Universidade Federal de São Paulo, UNIFESP, Rua Botucatu, 862 Vila Clementino, 04023-062 São Paulo, SP, Brasil
2Departamento de Morfologia e Fisiologia, Faculdade de Medicina do ABC, Avenida Lauro Gomes, 2000 Vila Sacadura, 09060-650 São Paulo, SP, Brasil
3Departamento de Biociências, Universidade Federal de São Paulo, UNIFESP, Avenida Ana Costa, 95 Vila Mathias, 11060-001 Santos, SP, Brasil
4Faculdade de Educação Física, Universidade Federal de Uberlândia, UFMG, Avenida Benjamin Constant, 1286 Bairro Aparecida, 38400-078 Uberlândia, MG, Brasil

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ABSTRACT

Several forms of experimental evidence gathered in the last 37 years have unequivocally established that the medulla oblongata harbors the main neural circuits responsible for generating and maintaining the sympathetic vasomotor tone and arterial blood pressure. The medulla oblongata also contains the main site integrating signals arising from high- and low-pressure baroreceptors and chemoreceptors afferents. Dysfunctions of this circuitry are a common feature of many pathological conditions and may be the core of cardiovascular diseases including arterial hypertension.

Our current views of this circuitry derive mainly from evidence gathered in the last 35 years, especially those from the studies of Pedro Guertzenstein and his colleagues. The fact that Guertzenstein was a former medical student at UNIFESP who became Professor of Physiology at UNIFESP later, and his colleagues. In this review, we have summarized the main findings as well as our collaboration to a further understanding of the ventrolateral medulla and the control of arterial blood pressure under normal and pathological conditions.

Key words: hypertension, baroreceptor reflexes, vasomotor nuclei, sympathetic nerve activity, arterial pressure.

INTRODUCTION

Several forms of experimental evidence gathered in the last 37 years have unequivocally established that the medulla oblongata harbors the main neural circuits responsible for the regulation of arterial blood pressure. Within this region, discrete groups of neurons act to generate and maintain the sympathetic vasomotor tone and arterial blood pressure. The medulla oblongata also contains the main site integrating signals arising from high- and low-pressure baroreceptors and chemoreceptors afferents. Dysfunctions of this circuitry are a common feature of many pathological conditions and may be the core of cardiovascular diseases including arterial hypertension.

Our current views of this circuitry derive mainly from evidence gathered in the last 35 years, especially those from the studies of Pedro Guertzenstein and his colleagues. The fact that Guertzenstein was a former medical student at UNIFESP who became Professor of the Department of Physiology later, and that many of us were able to contribute to the current view of cardiovascular regulation is a reason of pride and joy in the year in which UNIFESP commemorates its 75 years of foundation.

*Member Academia Brasileira de Ciências
Correspondence to: Dr. Sergio L. Cravo
E-mail: slcravo@fcr.epm.br
THE EARLY YEARS

The quest for the localization of the area responsible for the maintenance of vasomotor tone remounts to the XIX century. However, almost a century elapsed between C. Dittmar’s first attempts in Carl Ludwig’s laboratories in Leipzig (1873) and the initial experiments performed by P. Guertzenstein in William Feldberg’s laboratories in London (1971). Although Dittmar’s experiments led him to the conclusion that there was a vasomotor center localized in the lower half segment of the medulla, that is the ventral portion, his observations did not allow him to differentiate between the tonic and the reflexogenic areas of the supposed center. Furthermore, his anatomical definition was still preliminary since it included a fairly large area. Based on the results obtained by Owsjannikow (1871) and Dittmar (1873) in cats, the vasomotor center could be located anywhere in an area of 4 mm of length along the cranio-caudal axis (starting at the obex) and comprehending another 4 mm of ventral tissue in the mediolateral plane.

Nearly one hundred years later, in March 1970, Pedro G. Guertzenstein, at that time a young scientist from Brazil, arrived in Feldberg’s laboratories for a post-doctoral fellowship, staying there for three years. According to Feldberg’s own words: “We, that is, Guertzenstein and myself, stumbled on the ventral surface of the brain as late as 1972. ‘Our story’ began with a simple experiment, with a fall in arterial blood pressure following the injection of a few milligrammes of pentobarbitone sodium (Nembutal) into a lateral cerebral ventricle.” (Feldberg 1982).

During those three years, alone or in collaboration with many colleagues, Guertzenstein produced the impressive number of four communications to the Physiological Society (Guertzenstein, January 1971, Feldberg and Guertzenstein, January 1972, Guertzenstein, April 1972, Guertzenstein and Silver, June 1973) and five full papers published either in the Journal of Physiology or in the British Journal of Pharmacology (Feldberg and Guertzenstein 1972, Guertzenstein 1973, Bousquet and Guertzenstein 1973, Guertzenstein and Silver 1974, Edery and Guertzenstein 1974). Together with papers published much later after his return to Brazil, and including some developed during his last years at UNIFESP, these papers established the foundations of our current view of the ventrolateral medullary vasomotor nuclei and their role in the arterial blood pressure regulation. Since their publication, they were cited an average of 33 times/year, in a total amount of almost 1300 citations.

Out of these, the far most quoted and recognized as a classical paper is the one he published with the collaboration of Ann Silver (Guertzenstein and Silver 1974). In this paper they defined, for the very first time, the precise location of what is clearly recognized, until nowadays, as the rostroventrolateral medulla (RVLM), one and so far the most important source of tonic excitation to the sympathetic preganglionic neurons in the intermediolateral cell column of the spinal cord. Their results demonstrated unequivocally that, after a bilateral electrolytic destruction of a small area, not larger than 1 mm$^2$ in the ventrolateral medulla, the blood pressure was no longer maintained and remained low for at least 6 hours. In Guertzenstein’s own words: “Bilateral electrolytic destruction of the glycine sensitive area (GSA) produced a fall in arterial blood pressure to levels similar to those usually obtained in acute spinal animals, without signs of recovery for at least 6 h, which was the longest period of time the animals were observed for.” (Campos and Guertzenstein 1989).

The question about the role of the central nervous system in keeping arterial blood pressure levels was moved from where to how? The revolution in the way we thought about blood pressure regulation was shortly and precisely expressed by Feldberg (1982): “For a century the structures responsible for maintaining arterial blood pressure were thought to lie near the dorsal surface of the brain stem, on the floor of the fourth ventricle. Later, they were thought to be distributed more or less throughout the entire substance of the brain stem. Now we suggest that blood pressure may be maintained by the action of nerve cells located in a small bilateral region near its ventral surface”.

Shortly after, in 1976, Feldberg and Guertzenstein published another fundamental paper showing the existence of a different area, caudal to the one already described, on which topic application of nicotine produced a marked fall in blood pressure due to the inhibition of the vasoconstrictor tone. Assuming that nicotine was acting as an excitatory drug, they proposed: “With the
evidence so far available... there are at least two separate regions... a more rostral and a more caudal one” and also “yet the action itself is probably an excitatory one exert on inhibitory neurons that form connexions with the vasomotor pathway”. With these suggestions they had described what we would come to know as the caudal ventrolateral medulla (CVLM), and advanced the main properties of this region: its vasodepressor role through tonic and reflex inhibition of RVLM. A further characterization of this area in regulating cardiovascular functions, and particularly modulating cardiovascular reflexes, was developed after Guertzenstein’s return to Brazil, and was pioneerly presented in a communication followed by a full paper (Guertzenstein and Lopes 1980, 1984). The route for the understanding of the CVLM and its implications on the regulation of sympathetic tone and on cardiovascular reflexes was fully open and ready to be understood.

Many years later, based on a rather puzzling set of experimental observations, Guertzenstein and Feldberg went on to propose the existence of a yet third vasomotor area in the ventrolateral medulla. Once more their vision was far ahead of their time. A further development in the characterization of the area they foresaw took another 10 years. This was also his final enterprise because of his premature death in 1994. However, in his last papers, he and his fellows in the Department of Physiology at UNIFESP were able to show that the third area, the caudal pressor area (CPA), contains cells with a tonic pressor activity that contributed to the maintenance of baseline levels of arterial pressure and furthermore, that the CPA-induced cardiovascular responses were mediated by CVLM, with the involvement of both glutamatergic and GABAergic synapses (Possas et al. 1994, Campos et al. 1994).

Figure 1 contains a schematic representation of our current view of the main vasomotor nuclei in the ventrolateral medulla as derived from the work of Guertzenstein and colleagues, and has been confirmed in literally thousands of papers published in the last 30 years. In this review, we summarized the main findings and our collaboration to the development in the knowledge of the ventrolateral medulla and the tonic and reflex regulation of the arterial blood pressure.

The early studies performed in cats by Guertzenstein and Feldberg were a landmark in our understanding of the mechanism by which the sympathetic vasomotor tone is generated by the central nervous system. There is now considerable evidence that a restricted group of specialized reticulospinal neurons located in the RVLM is crucial to maintain the sympathetic vasomotor tone in different species. The major characteristics of the RVLM neurons include: direct monosynaptic excitatory connections to identified pre-ganglionic sympathetic neurons of the spinal cord, tonic activity and baroreceptor sensitivity.

The RVLM receives inputs from a number of different nuclei in the brain and also sends projections to many other regions involved in the cardiovascular, respiratory and hormonal control. A number of reciprocal innervations between the RVLM and other brain nuclei strongly suggest that the RVLM is not only an important region involved in the maintenance of the tonic sympathetic vasomotor tone. It might also be an integrative center controlling the cardiovascular functions, processing the information from the peripheral nerves (baroreceptor and chemoreceptor reflexes) and from other nuclei acting as a key region to maintain cardiovascular homeostasis.

The RVLM neurons have been extensively studied using different approaches. First, they were identified by topical application of drugs (glycine or GABA) in the ventral surface of the brainstem by Guertzenstein and colleagues. When applied to the rostral part of the ventrolateral medulla in cats, such amino acids caused a large fall in blood pressure and cardiac output in the region denominated as the glycine sensitive area, now denominated the RVLM (Guertzenstein and Silver 1974, Campos and Guertzenstein 1989). Subsequently, the anatomical location of the RVLM was defined in the rat and in the rabbit using microinjections of amino acids directly into the ventrolateral medulla parenchyma (Ross et al. 1984). A more precise localization of the RVLM neurons was achieved, the cardiovascular neurons were then localized ventrally to the rostral part of the nucleus ambiguous (Dampney 1994). The precise location of the RVLM region allowed the study of these neurons with more refined and improved techniques including single
Fig. 1 – A: The current view of the role of the ventrolateral medulla and cardiovascular regulation: tonic sympathetic activity to the heart, resistance vessels and adrenal medulla derive from preganglionic sympathetic neurons (SPNs) located in the intermediolateral cell column. SPNs are tonically excited by direct bulbospinal neurons located in the rostral ventrolateral medulla (RVLM). Activity of RVLM neurons is regulated by inhibitory afferents located in the caudal ventrolateral medulla (CVLM) and by activity of neurons in caudal pressor area (CPA). The nucleus tractus solitarius (NTS) is the primary site receiving afferents from high and low pressure baroceptors and chemoreceptors. From the NTS this information is transferred to the VLM. Arterial baroreceptors reflexes are mediated by the inhibitory CVLM-RVLM pathway. From CVL ascending efferents project to several nuclei involved in water and salt intake and cardiovascular control, e.g., the Median Preoptic Nucleus (MePO), the paraventricular nucleus of hypothalamus (PVH) and the supraoptic nucleus (SON). B: Schematic representation of three coronal sections of the rat medulla oblongata at the levels of the RVLM, CVLM and CPA, respectively. The hatched areas represent the areas from which characteristic cardiovascular responses can be evoked. C: Diagram representing the rat’s ventral medullary surface showing the localizations (from rostral to caudal) of the RVLM, CVLM and CPA (Modified from Cravo et al. 2006).

unit electrophysiology, immunohistochemistry and cellular molecular biology techniques.

The RVLM neurons have been studied using extracellular recording (Brown and Guyenet 1984, 1985; Morrison et al. 1988, Campos and McAllen 1999) and intracellular recordings (Lipski et al. 1996). However, despite the large number of studies on the RVLM and its recognized importance on the sympathetic vasomotor tone generation, the basis for the tonic ongoing activity of these neurons is not yet fully understood.
At least, under specific experimental conditions in vivo, fast excitatory synaptic inputs (EPSPs) appear to drive the RVLM spiking activity. The ongoing activity of these neurons resulted of synaptic inputs, with individual action potentials usually preceded by identifiable fast EPSPs (Lipski et al. 1996). These findings are in agreement with the network hypothesis of the generation of sympathetic vasomotor tone proposed by Barman and Gebber (1987). The hypothesis implies that the activity of premotor neurons in vivo is dependent on excitatory inputs from other brainstem nuclei. The question is: where are the sources of the synaptic drive to the RVLM?

There is a large body of evidence showing a number of nuclei in the brain and peripheral nerves from which synaptic excitation of RVLM neurons can be achieved, following electrical or chemical stimulation (Sun and Guyenet 1986, Cechetto and Chen 1992). Although a number of brain stem regions, when activated, cause sympathetic activation via the RVLM, few regions may provide a tonic excitatory drive to the RVLM neurons to support their activity. So far, some regions in the brain stem have been identified that may provide a tonic drive to the RVLM presympathetic neurons. Among these regions are:

1) the lateral tegmental field (LTF) in the dorsal formation of the medulla oblongata (Barman and Gebber 1987),
2) the pontine reticular formation (Hayes and Weaver 1992) and,
3) the caudal pressor area (CPA) in the caudal end of the ventrolateral medulla (Campos and McAllen 1999).

The LTF has a sympathetic-related activity and contains neurons that respond to the baroreceptor reflex, with an increase or a decrease in their activity in response to baroreceptor activation. The barosensitive LTF neurons send projections to the RVLM and are probably one source of excitatory inputs to the region. Furthermore, blockade of N-methyl-D-aspartate (NMDA) receptors in the LTF abolished baroreceptor reflex control of sympathetic activity (Barman and Gebber 1987). In a recent study in cats, the same group showed that a blockade of non-NMDA receptors in the LTF significantly attenuated the reflex increase in cardiac and vertebral sympathetic nerve activity in response to electrical stimulation of vagal afferents or by activation of arterial chemoreceptors (Orer et al. 2004). On the other hand, the reflex sympathetic activation, in response to electrical stimulation of the sciatic or trigeminal nerve, was not affected by previous glutamatergic blockade within the LTF. These data suggest an important and specific role of the LTF controlling the sympathetic reflex pathways. However, the role of this region in supporting the vasomotor tone needs to be clarified. Hitherto the evidence that blood pressure falls when the LTF cell bodies are inactivated is lacking. Furthermore, the anatomical localization and the role of LTF neurons in rats are not very well defined, and the possibility that there is some homology between the LTF in the cat and the CVLM in rats cannot be ruled out and needs to be clarified.

A second region that can support the RVLM ongoing activity is the pontine reticular formation. Hayes and Weaver (1992) found that glycine microinjection into a diffuse region of the pontine reticular formation caused a decrease in blood pressure in anesthetized rats. However, the full meaning of this interaction with the vasomotor tone needs to be more thoroughly studied.

Finally, the CPA is probably an important source to maintain the RVLM activity. The CPA was discovered in cats by Feldberg and Guertzenstein (1986) and in rats by Gordon and McCann (1988). In anesthetized rats, bilateral inhibition of the CPA by GABA or glycine decreased the blood pressure by 30–40 mmHg (Campos et al. 1994). In rabbits, CPA inhibition caused a similar decrease in the arterial pressure and almost abolished the renal sympathetic nerve activity (Dampney et al. 2003). The decrease in the arterial pressure is probably mediated by a decrease in RVLM activity. It was shown that the RVLM sympathetic premotor neurons were inhibited on an average of 40% during a unilateral or bilateral microinjection of glycine into the CPA (Campos and McAllen 1999). Taken together, these data suggest that an important fraction of resting activity of sympathetic premotor neurons of the RVLM depends on a synaptic drive from the CPA. However, the exact physiological role of the CPA is not yet known. Furthermore, there is no information on what drives the CPA neurons and what kind of neurons exists within this region.

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Concluding, despite the large number of studies, understanding how RVLM neurons work to maintain the sympathetic vasomotor drive in non-anesthetized animals remains a challenge, and the characteristics of depolarization and the membrane properties of the RVLM neurons in intact and conscious animals are not fully understood.

There are two major indications that the RVLM is involved in the long-term control of sympathetic activity and blood pressure:

1) pharmacological evidence indicates that the RVLM is the major site of action of centrally antihypertensive agents such as clonidine and moxonidine (Ernsberger et al. 1987), and

2) experimental fulminating neurogenic hypertension induced through lesion of the CNS at sites such as the nucleus of the solitary tract or the caudal ventrolateral medulla occurred as a consequence of RVLM disinhibition (Doba and Reis 1973, Blessing and Reis 1982). Recently, this hypothesis was supported by a new evidence showing that changes in neurotransmission within the RVLM are related to acute or chronic hypertension.

Electrophysiological studies in spontaneously hypertensive rats (SHRs) have confirmed increases in the firing rates of RVLM barosensitive neurons containing excitatory amino acids and terminating in the intermediolateral cell column of the spinal cord, where the preganglionic sympathetic neurons SPNs are located (Chalmers et al. 1992, Chan et al. 1991). We can hypothesize that an increase in excitatory neurotransmission in the RVLM, glutamatergic activity in particular, is in part responsible for the sympathetic activation seen in experimental hypertension.

In an experimental model of renovascular (two-kidney, one-clip Goldblatt) hypertension in rats, bilateral blockade of RVLM glutamatergic synapses by micro-injection of kynurenic acid, a broad-spectrum glutamate receptor antagonist, decreased blood pressure to a degree similar to that caused by ganglionic blockade. In addition, the blockade of excitatory amino acid input to the RVLM did not alter the blood pressure in normal animals (Bergamaschi et al. 1995). These findings suggest that, in the RVLM, glutamate plays a tonic role in renovascular hypertensive animals, but not in normotensive animals. Figure 2 shows a representative recording of blood pressure and renal sympathetic activity during glutamatergic synapses blockade into the RVLM of a renovascular hypertensive rat.

The mechanisms by which glutamate is activated in the RVLM during renovascular hypertension are not fully understood. One possibility is that a high level of circulating angiotensin II (Ang II) excites RVLM neurons directly or indirectly, causing sympathoexcitation. It has been documented that direct injection of Ang II into the RVLM caused blood pressure to increase in anesthetized animals (Allen et al. 1988, Andreatta et al. 1988, Sakai and Dampney 1990). In addition, introduction of an Ang II antagonist into the RVLM has been shown to produce significant drops in the blood pressure and sympathetic activity (Sakai and Dampney 1990, Averill et al. 1994, Ito and Sved 1996). These findings suggest that endogenous Ang II causes tonic excitation of RVLM neurons, and this may be relevant to sympathetic activation in hypertension.

In renovascular hypertensive rats, as well as in SHRs (Ito et al. 2000), Dahl salt-sensitive rats (Ito et al. 2001) and rats used in an experimental model of pulsatile compression of the RVLM (Morimoto et al. 1999), the blockade of excitatory amino acid receptors in the RVLM elicited profound decreases in the arterial pressure. Our hypothesis is that the increased glutamatergic activity in the RVLM seen in hypertension is in part mediated by local or circulating Ang II.

This hypothesis is supported by studies in renovascular hypertensive rats, in which the depressor effect of glutamate blockade in the RVLM was not observed in animals pretreated with a low dose of an Ang II enzyme converter inhibitor (captopril), suggesting that the glutamatergic activation of the RVLM in this model depended on high levels of circulating Ang II (Carvalho et al. 2003).

Other evidence supporting an increase of glutamatergic inputs to the RVLM, in response to Ang II, has been provided by studies using microdialysis, in which intravenous infusion of Ang II at a very low rate over a period of several hours caused a significant increase in glutamate release in the RVLM (Katahira et al. 1994, Moriguchi et al. 1994). In addition, intravenous admin-
administration of an angiotensin-converting enzyme inhibitor caused a decrease in glutamate release in the RVLM (Katahira et al. 1994, Moriguchi et al. 1994). All these data support the idea that there is an interaction between circulating Ang II and glutamatergic drive to the RVLM. However, the cellular mechanisms by which Ang II causes an increase in glutamatergic actions in the RVLM have yet to be clarified.

Since circulating Ang II does not have direct access to the RVLM, the excitatory effect of Ang II is likely to be derived from angiotensinergic neural inputs or from paracrine secretion within the ventrolateral medulla (Van et al. 1980, Fink 1997). Central AT1 receptor-mediated Ang II activity is mediated by facilitation of excitatory transmission, not only via glutamate but also via catecholamine release (Ferguson et al. 2001), substance P (Paton et al. 2001) and, paradoxically, enhanced inhibitory GABAergic neurotransmission (Paton and Kasparovo 1999).

A model postulated to explain the sympathoexcitation that Ang II causes via the RVLM was proposed years ago by Joy and Lowe (1970). According to their hypothesis, the area postrema (AP) is an important locus at which circulating Ang II modulates the sympathetic vasmotor tone originating in the ventrolateral medulla. Ablation of the AP significantly inhibited the increase in the blood pressure associated with chronic intravenous infusion of Ang II in rats (Fink et al. 1987). A direct or indirect neural pathway from the AP to the RVLM might be involved in the increased excitatory neurotransmission mediated by circulating Ang II. In addition, local paracrine secretion of Ang II in the ventrolateral medulla may participate in this mechanism.

It is interesting that, in other models of sympathetic activation such as acute or chronic water deprivation, there is an increase in excitatory amino acid drive in the RVLM that is caused specifically by an increased osmolarity rather than by a decreased blood volume (Brooks et al. 2004).

We can speculate that the sympathoexcitation may be in part a consequence of an increase in the glutamatergic actions within the RVLM, not only in experimental models of hypertension, but also under other conditions associated with increased sympathetic drive. As proposed by Lipski (Lipski et al. 1996), it is possible that, under normal conditions, RVLM premotor neuron activity is determined by the balance between excitatory and inhibitory synaptic inputs, including amino acid excita-
tery mechanisms. However, under special conditions, such as hypertension, an imbalance between these inputs, resulting in an increased excitatory activity within the RVLM, may enhance the tonic activity of RVLM neurons.

**THE CAUDAL VENTROLATERAL MEDULLA (CVLM)**

The term caudal ventrolateral medulla (CVLM) was initially proposed by Reis and coworkers (Ross et al. 1984) and defined essentially in analogy with the rostral ventrolateral medulla. The CVLM, as functionally defined in the rat, is certainly the equivalent of the nicotine area described by Feldberg and Guertzenstein in cats (Feldberg and Guertzenstein 1976). However, while the RVLM can be functionally and anatomically defined in a rather discrete area, the CVLM is a much larger area whose boundaries are not yet precisely established. In the rat, CVLM neurons are spread in the rostrocaudal axis from the vicinity of the RVLM to the spinal-medullary junction (Ross et al. 1985, Ruggiero et al. 1994). The CVLM is also a functionally heterogeneous area. It contains at least two separate systems that are distinct in their connections and function. These circuits are involved in the control of peripheral resistance and the volume and composition of the extracellular compartment.

The CVLM circuitry for controlling peripheral resistance is represented by a group of sympatoinhibitory neurons scattered around the periambigual area whose function is essential in cardiovascular regulation. CVLM stimulation produces marked hypotension and bradycardia due to a reduction in the sympathetic vasoconstrictor drive (Blessing and Reis 1982, Cravo et al. 1991). Anatomical and functional evidence indicate that CVLM sympatoinhibition is mediated through the inhibition of RVLM neurons (Granata et al. 1986, Willete et al. 1987). Retrograde tracing studies demonstrated that CVLM neurons do not contact SPNs in the intermediolateral cell column, but sent dense projections to the RVLM (Ross et al. 1985). There, putative GABAergic CVLM neurons form inhibitory synapses with bulbospinal RVLM neurons (Aicher et al. 1996), providing a tonic GABAergic inhibition. When the CVLM is lesioned or functionally inactivated, there are marked increases in sympathetic nerve activity and neurogenic hypertension (Blessing et al. 1982, Cravo et al. 1991). This hypertension can reach levels capable of producing ventricular failure and death due to acute ventricular failure and pulmonary edema.

Tracing studies also indicate that the CVLM receives numerous projections from the NTS areas receiving primary baroreceptor afferent fibers (Ross et al. 1985). It is now accepted that the CVLM contains an essential interneuron of the baroreceptor reflex arch. Inactivation of the CVLM abolished baroreceptor adjustments (Cravo et al. 1991, Granata et al. 1986). Extracellular recordings have identified within the CVLM neurons with all characteristics of a baroreceptor interneuron: orthodromical activation by electrical stimulation of baroreceptor afferents, NTS stimulation or increases in the arterial blood pressure and direct projections to RVLM (Jeske et al. 1993). From the work of Feldberg, Guertzenstein and Lopes, our recognition of the CVLM reflex and tonic sympathoinhibition has come a long way. However, a key aspect of CVLM functions remains largely unknown: similarly to the rostral medulla, neurons of the CVLM exhibit tonic activity which is under a tonic GABAergic inhibitory control. The origins of both the tonic activity and the GABAergic afferents are unidentified. Results obtained with selective microinjection of glutamatergic and GABAergic antagonists suggested that CVLM tonic activity is maintained through a combination of tonic excitatory and inhibitory afferents whose source, however, remains largely unknown (Guyenet et al. 1987, Campos Jr. et al. 1994).

Besides its role in the peripheral resistance regulation, the CVLM neurons are also important for body fluid homeostasis. In the beginning of the 1960s, Dahlström and Fuxe (1964) showed the presence of catecholaminergic neurons in this area (A1 noradrenergic neurons). Initially it was thought that A1 neurons were part of the CVLM sympathoinhibitory group (Granata et al. 1986), but later it became clear that these cells were GABAergic and located medially to the A1 neurons. Now it is well accepted that A1 neurons are central elements in regulating the volume and composition of the extracellular compartment (Pedrinò et al. 2008, 2006, Howe et al. 2004, Buller et al. 1999, Hochstenbach and Ciriello 1995, Smith et al. 1995, Head et al. 1987).

Neuroanatomical studies have shown that A1 noradrenergic neurons receive projections from arterial baro-
Cardiovascular control by the ventrolateral medulla

Receptors as well as from vagal cardiopulmonary volume receptors (Li et al. 1992, Day et al. 1992, Day and Sibbald 1990). These neurons also are reciprocally connected with hypothalamic regions known for their involvement in neuroendocrine, hydroelectrolytic, and cardiovascular regulation, including the median preoptic nucleus (MnPO), the subfornical organ (SFO), the paraventricular nucleus (PVN), and the supraoptic nucleus (SON, Tanaka et al. 1997, Tucker et al. 1987, Saper et al. 1983).

Results obtained in studies developed in the last 10 years in our group demonstrated that A1 neurons are part of central pathways involved in cardiovascular responses induced by acute changes of volume and composition of the extracellular compartment. In intact anesthetized rats, isotonic volume expansion or hypernatremia induced a sustained renal vasodilation. The observed renal vasodilation is due to a reduction of the renal sympathetic nerve activity and the release of vasoactive peptides. Denervation of the baroreceptors or electrolytic lesions of the anterioventral wall of the third ventricle of the brain (AV3V) abolished renal vasodilation (Pedrino et al. 2005, Colombari et al. 2000, Colombari and Cravo 1999). Blockade of the adrenergic transmission in the AV3V reduced ANP release and renal vasodilation induced by changes in circulating volume (Antunes-Rodrigues et al. 1993, Pedrino and Cravo 2006 Abstract). Taken together, these findings suggest that the catecholaminergic innervations, originating mainly from A1 neurons, represent a necessary neural pathway involved in responses to acute changes in the volume or composition of the extracellular compartment.

A functional relationship between A1 neurons and cardiovascular and humoral responses to acute reductions in central blood volume was demonstrated in previous studies (Buller et al. 1999, Smith et al. 1995, Head et al. 1987). Buller et al. (1999) demonstrated that the lesion of the region where A1 noradrenergic neurons are located reduced the number of Fos-positive neurosecretory vasopressin cells in the SON and PVN induced by hypotensive hemorrhage. Similarly, other studies demonstrated that lesions of these neurons reduced the vasopressin secretion induced by a decreased circulating volume (Smith et al. 1995, Head et al. 1987). Despite the abundance of evidence provided by studies to support this important role of A1 noradrenergic neurons in acute reductions of blood volume, little is known regarding the role of these neurons during hypernatremia or volume expansion. Previous studies employing early gene expression have demonstrated that subcutaneous, intraperitoneal or intravenous administration of hypertonic saline or isotonic volume expansion increases Fos and c-Fos mRNA expression in the A1 noradrenergic neurons (Howe et al. 2004, Hochstenbach and Ciriello 1995). Moreover, in recent studies, we demonstrated that renal vasodilation and sympathoinhibition induced by increases in plasma sodium concentration was blunted in animals submitted to specific lesion of A1 noradrenergic neurons (Pedrino et al. 2006). These studies are the initial observations that A1 noradrenergic neurons are involved on sympathetic and cardiovascular changes induced by hypernatremia.

Overall, these recent evidence support the idea that A1 neurons in the CVLM are activated upon stimulation of peripheral baroreceptor and cardiopulmonary afferents, engaging efferent pathways to AV3V and PVN that regulate the endocrine and autonomic responses relevant to body fluid homeostasis and cardiovascular regulation. Since A1 neurons are important to sympathetic and peripheral resistance response to changes of circulating volume, the irregular function of these neurons results in an inefficient control of the renal sympathetic activity which could contribute to the pathophysiology of hypertension, congestive heart failure and cirrhosis.

The nucleus of the solitary tract (NTS): old and new pathways toward the ventral surface of the medulla

Different evidence has shown that the NTS constitutes the primary site of integration of different visceral sensory afferents. The intermediate and commissural portions of the NTS are mostly known as the “cardiovascular NTS”. The intermediate NTS receives mainly carotid and aortic baroreceptor afferents, while the commissural NTS is the site of termination of arterial chemoreceptors and also aortic baroreceptor fibers (Cottle 1964, Crill and Reis 1980, Miura and Reis 1969, 1972, Spyer 1981, Finley and Katz 1992, Ciriello et al. 1994, Colombari et al. 1996). Information arriving from these afferents...
into the NTS, which in turn sends efferent projections to different areas involved in cardiovascular regulation. The ventrolateral medullary surface is a main site of these efferents.

Studies have shown that NTS efferent terminals synapse onto neurons in the CVLM that project to the RVLM (Aicher et al. 1996). Although neuroanatomical and functional evidence suggest that the information of arterial baroreceptors are integrated in the classic circuit NTS-CVLM-RVLM, different findings (Willete et al. 1984a, b) have shown that the stimulation of the aortic depressor nerve elicited a pressor response in rats with inhibition of CVLM neurons. Indeed, this apparently paradox effect that differs from the expected hypotension and bradycardia evoked by the aortic baroreceptor stimulation suggested that the pressor response would be dependent on a direct pathway from NTS to the RVLM. Urbanski and Sapru (1988a, b) have shown that a microinjection of L-glutamate into the NTS in anesthetized rats that had the ipsilateral CVLM inhibited with muscimol or lidocaine produced a pressor response. This effect was the opposite from the expected response in anesthetized rats with an intact CVLM. Immunohistochemical studies showed the existence of projections from the NTS to the RVLM (Ross et al. 1985). In a different approach, NTS glutamatergic stimulation in a non-anesthetized condition consistently produced increases in arterial pressure. However, in commissural NTS-lesioned rats, such pressure response was turn in a fall in arterial pressure as demonstrated in anesthetized animals (Colombari et al. 1996). Taken together, these data support two hypotheses: a) in addition to the sympatoinhibitory projections from NTS to the ventral surface, the NTS drives sympatoexcitatory pathways as well; b) this sympatoexcitatory responses could be integrated via a direct NTS-RVLM pathway.

Further studies have shown that the pressor responses induced by stimulation of arterial chemoreceptors with N2 were not affected by CVLM inhibition neurons or the blockade of glutamatergic transmission in the CVLM (Koshiya et al. 1993). Nevertheless, glutamatergic blockade in the RVLM abolished these pressor responses. These results suggest that the sympatoexcitatory component of the chemoreceptor reflex response depends on a possible direct glutamatergic pathway from the NTS to the RVLM (Koshiya et al. 1993). Electrophysiological evidence has also shown that chemosensitive neurons in the commissural subnucleus of the NTS are antidromically activated by the RVLM, which suggested that commissural NTS neurons arborize in the RVL (Koshiya and Guyenet 1996). These findings support the idea that not only the intermediate but also the commissural subnucleus of the NTS would project directly to the RVLM.

Although the commissural subnucleus of the NTS is specially known for integrating the chemoreceptor afferent information, it has been shown that lesions of this subnucleus of the NTS abolished the hypertensive response evoked by the aortic baroreceptor denervation (Sato et al. 1999). Electrolytic lesions of the commissural NTS also elicited a marked fall in arterial pressure in spontaneously hypertensive rats (SHR), but not in normotensive Wistar-Kyoto rats (Sato et al. 2001). These lesions abolished the pressor response of the chemoreceptor reflex and attenuated the sympathoexcitatory component of the baroreceptor reflex in SHR (Sato et al. 2001, Colombari et al. 2001). Inhibitions of the commissural NTS neurons have also reduced the splanchnic sympathetic nerve activity and, consequently, the arterial pressure in SHR, but not in normotensive rats (Sato et al. 2002). Indeed, the integrity of the commissural NTS seems to be essential for the development of the hypertensive response in aortic denervated rats or for the maintenance of the high blood pressure in SHR, which suggest a possible tonic activity of these neurons for the hypertensive condition. In addition, the microinjection of substance P into the commissural NTS of juvenile SHR reduced the thoracic sympathetic trunk nerve activity and induced vasodilatation, but not in normotensive WKY rats. Hence, a possible neurotransmitter which might be up-regulated in this hypertension model would be the substance P (AP Abdala et al., unpublished data). Other studies have shown that the blockade of excitatory amino acid receptors in the RVLM reduced arterial pressure in SHR with little effect in normotensive rats (Ito and Sved 1997). Taken together, these evidences suggest that the commissural NTS is likely to be an important source of excitation to RVLM neurons and, therefore, may constitute a neural pathway which can be altered and possibly tonical-
ly active under pathophysiological conditions such as hypertension.

Although either the intermediate or the commissural NTS have been shown as a possible source for a direct excitation of the RVLM, whereas the CVLM has been known as a source of RVLM inhibition, new findings suggest changes in this view of the brain stem network. Moreira et al. (2005) showed that the blockade of the glutamatergic receptors in the NTS or the inhibition of the CVLM increased arterial pressure and heart rate as previously demonstrated (Guyenet et al. 1987, Willette et al. 1987, Dampney 1994, Ito and Sved 1997). However, a new finding of this study was the demonstration that, after the simultaneous blockade of the glutamatergic receptors of the NTS and the inhibition of the CVLM, blood pressure and mesenteric and hindquarter vascular resistances were reduced below baseline levels. These findings suggest the existence of an important pressor mechanism arising from the NTS, and also that an excitatory pathway from the CVLM to the RVLM is likely to be involved in the control of vascular resistance and arterial pressure.

CONCLUDING REMARKS

The work of Guertzenstein and colleagues has defined what today we recognize as the main circuitry in the generation of the vasomotor sympathetic activity and blood pressure regulation. His pioneering work endowed our comprehension of blood pressure regulation. Several main questions remain open and we hope to continue contributing to their answers. This review is dedicated to him, a distinguished scientist and a dearly missed friend.

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RESUMO

Numerosas formas de evidência experimental obtidas nos últimos 37 anos demonstraram inequivocamente que a medula oblongata contém os principais circuitos responsáveis pela geração e manutenção do tônus vasomotor e a regulação da pressão arterial. A visão atual que possuímos destes circuitos deriva em grande parte dos estudos de Pedro Guertzenstein, um estudante e mais tarde Professor de Fisiologia da UNIFESP e seus colaboradores. Nesta revisão nós sumarizamos os seus principais resultados assim como a nossa colaboração para uma melhor compreensão da regulação da pressão arterial em condições normais e patológicas.

REFERENCES


CARdiovascular Control By THE VENTROLATERAL Medulla


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