Ion homeostasis in rhythmogenesis: The interplay between neurons and astroglia.

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Abstract

Proper function of all excitable cells depends on ions homeostasis. Nowhere is this more critical than in the brain where the extracellular concentration of some ions determines neurons firing pattern and ability to encode information. Several neuronal functions depend on the ability of neurons to change their firing pattern to a rhythmic bursting pattern, whereas, in some circuits, rhythmic firing is, on the contrary, associated to pathologies like epilepsy or Parkinson's disease. In this review, we focus on the four main ions known to fluctuate during rhythmic firing: calcium, potassium, sodium and chloride. We discuss the synergistic interactions between these elements in order to promote an oscillatory activity. We also review evidence supporting an important role for astrocytes in the homeostasis of each of these ions and describe mechanisms by which astrocytes may regulate neuronal firing by altering their extracellular concentrations. A particular emphasis is put on the mechanisms underlying rhythmogenesis in the circuit forming the central pattern generator (CPG) for mastication and other CPG systems. Finally, we discuss how an impairment in the ability of glial cells to maintain such homeostasis may result in pathologies like epilepsy and the Parkinson's disease.

Keywords

Astrocytes, Rhythmogenesis, Homeostasis, Calcium, Potassium

I- INTRODUCTION

Proper neuronal function depends critically on the ability of neurons to encode information by changing their firing pattern. Whether a neuron unfailingly transmits spike trains or generates an oscillatory burst firing in response to a given incoming input will bestow different encodings to the conveyed information, and result in totally distinct output signals that have different functional incidence. In several brain areas, neurons have been shown to display a function-related dual mode of firing alternating between a single spike firing mode and a rhythmic burst pattern (67, 92, 98, 111, 139). The role and necessity of such rhythmic neuronal discharges is obvious for some functions such as repetitive movements like mastication, locomotion or breathing. Indeed, for rhythmic movements to occur, repetitive motor output commands are required to produce the appropriate rhythmic contractions of the skeletal muscles involved in their execution. This implies a need for coordinated discharges by the central pattern generators (CPGs) to initiate and maintain the rhythmic output of motoneurons. Nevertheless, rhythmic activities are also observed in several areas of the brain where their function is not clearly related to a rhythmic behavior, but rather to functions such as the binding of sensory features in perception (51, 93), cognitive processing (91), information transfer between brain areas (131), learning and memory (138), sleep and consciousness (38) or motor coordination (79, 84). In fact, neuronal oscillatory activity appears to be a ubiquitous phenomenon in the brain with wide-ranging functions that occur along with normal brain processing (for review see (159)).

The mechanisms underlying the transition from single spike firing to rhythmic bursting are not fully understood yet. However, considering the critical importance of ion homeostasis for an optimal neural function, an obvious way to impact on the neuronal firing properties would be to modulate the ionic concentration in the extracellular compartment. Indeed, in several studies where the ions concentrations were measured directly in the tissue, variations in extracellular ions concentrations occurred prior or simultaneously to rhythmic neuronal discharges (3, 4, 19). Moreover, artificially decreasing the concentration of extracellular calcium, triggers rhythmic firing in masticatory and locomotor central pattern generator neurons (20, 149).

Astrocytes have been shown to modulate neuronal activity by a variety of mechanisms, but one of their most well established functions is to maintain ions homeostasis. Moreover, astrocytes are well positioned in term of their structural organisation relatively to neurons to monitor the extracellular space content. Thus, here we review the evidence suggesting that astrocytes may play

a determinant role in setting the firing mode of adjacent neurons through regulation of the extracellular ionic concentrations.

In the upcoming sections, we will discuss how the concentration of different ion species evolves during rhythmic oscillations. This review will put a special emphasis on potassium and calcium which are the most considered ions in studies interested in the generation and maintenance of rhythmic oscillations. Evidence supporting a role for glial cells in the generation of rhythmic firing in both healthy and pathological conditions will also be discussed.

II- RHYTHMIC OSCILLATIONS

2.1 Rhythmogenesis

Rhythmogenesis is the cellular process by which a rhythmic mode of firing or burst is generated at the cellular level. For the sake of clarity, we define bursts as recurrent clusters of action potentials occurring at short intervals interspike overriding а plateau-like depolarization and separated by relatively short periods of repolarization (see Figure 1A). Consequently, the main characteristic of the bursting pattern is its marked periodicity by contrast to the tonic spiking mode which consists in the firing of single action potentials that may be occasional or repetitive but that lacks the apparent recurrent on/off phasic features and that often occurs at lower frequencies than the firing overriding the plateaus that compose the bursts. Rhythmic clustering of neuronal discharge is sometimes also named neural oscillations and, under this terminology, refers as well to the oscillatory activity of individual neurons or to the largescale brain oscillations that result from recurrent activity in groups of neurons. These large-scale brain oscillations can be detected locally with field potentials recordings or measured outside the scalp by electroencephalograms (EEGs) and are more evocative of the state or degree of firing synchronicity in and between groups of neurons throughout the brain. This supposes that these oscillations, in some situations, may arise from a window of specific spatio-temporal configurations of the spiking activity of individual neurons within a group without an absolute requirement for rhythmic bursting in single cells activity. Nevertheless, numerous studies established irrefutable links between the large-scale brain oscillations and membrane potential fluctuations or rhythmic clustering of action potentials firing in individual neurons (78, 144). This suggests that, in many cases, these large-scale brain oscillations may depend on the generation of a rhythmic firing pattern at the cellular level. In fact, it has been postulated that bursting neurons might serve as pacemaker in network oscillations (29). Consequently, in this review, we will discuss and use the term rhythmogenesis to refer both to the generation of rhythmic patterns of action potentials at the cellular level and to the generation of the largescale brain oscillations.

2.2 Ionic basis of rhythmogenesis

The precise mechanisms leading to rhythmogenesis vary from one network to another and depend on an intricate interplay between synaptic and intrinsic properties in neuron populations. Many theories have been proposed to explain the generation of oscillatory activity including synaptic interactions and feedback connections within and between neuronal populations and their contribution to different known frequency bands (159). However, in this review, we will focus on intrinsic neuronal properties that rely on ionic conductances to generate rhythmic firing.

Some of these conductances constitute the driving forces that initiate the burst firing while others determine the plateaus duration and bursting frequency or are part of the regenerative feature that characterizes this mode of firing (57). The persistent sodium current (I_{NaP}) is one of the most common currents that drives bursting. Unlike the transient sodium current which appears at membrane potentials close to -50 mV (21), the I_{NaP} current becomes apparent at membrane potentials between -65 and -50 mV and peaks at membrane potentials between -40 and -35 mV, depending on the structure considered (21, 28, 100, 152). Some of the persistent current originates from the same pool of channels as the transient sodium current but with different gating modes (2) and in some other cases, as in the cerebellar Purkinje cells (156), it results from distinctive pools of channels. The massive entry of sodium ions into the cells causing the persistent sodium current supports depolarizing plateaus that are crucial for rhythmic bursting in some neurons. Evidence supporting a role for I_{NaP} in the generation of bursts came from computational simulations (68, 132) experimental recordings, from the preBötzinger complex (PBC) (36, 37), the main sensory nucleus of the trigeminal nerve (20, 152), the spinal cord (149), the cortex (23, 54, 110) or the hippocampus (72, 146). In these experiments, blockade of the I_{NaP} current prevented burst firing. The I_{NaP} current also contributes to large-scale oscillations like those in the rat's visual cortex, where its blockade significantly reduces the frequency of fast gamma oscillations (113).

Other drivers of burst firing include the low threshold voltage-activated calcium channels (LVA) and the calcium-activated non-selective cationic channels. The LVA calcium channels are found in a wide range of cell types including neurons and muscle fibers and are

generally activated by small voltage changes (6, 12) near the resting potentials of the membrane they are embedded in. The LVA calcium channels (mainly formed by the $Ca_{V}3$ family) mediate the T-current (I_{T}) responsible for the low-threshold calcium spikes responsible for burst firing and low-frequency oscillations in the thalamus and the cerebellum (for review see (66, 120)). At rest, the LVA calcium channels are in an inactivated state and require membrane hyperpolarisation to be de-inactivated allowing their subsequent activation upon membrane depolarisation. Activation of the LVA calcium channels by a subsequent depolarization causes a massive entry of calcium ions into the cell that generates sufficient depolarization to reach threshold and drive a train of Na⁺-dependent action potentials. Whole-cell extracellular recordings in the thalamus of mice knocked out for Ca_V3.2 and Ca_V3.3 clearly show the role that these channels play in regard to rhythmic bursting activity. In these experiments, periodic bursts were absent or significantly reduced in thalamic neurons for Ca_v3.3 KO mice (7) and simultaneously reduced and increased in the reticular and ventroposterior thalamic neurons respectively for mice devoid of Ca_V3.2 (88). The tonic discharges were unchanged (7). Ca_v1.3 calcium channels, members of the Ca_v1 family, could also contribute to burst firing. These channels, which are part of the L-type calcium channels, have been classified as high voltage activated (HVA) calcium channels, although they display, in reality, a very low voltage of activation (around -55 mV) (162). They are mostly known to contribute to pacemaking in cardiac atrial tissue and adrenal chromaffin cells, but may also play a role in neuronal bursting activity (89, 122).

The channels driving I_{CAN} are activated by the mobilization of intracellular calcium. These channels are seemingly part of the transient receptor potential (TRPM) family which includes TRPM4 and TRPM5. They are mainly permeable to potassium and sodium but very little to calcium (83, 154). I_{CAN} is responsible for prolonged plateaus (10, 164) seen during rhythmic bursting in several types of cells though they have no inherent voltage dependency. The contribution of these channels to bursting was confirmed with mathematical models (130) and in many experimental preparations including, the reticular thalamic neurons in the guineapig (10), the hypothalamic magnocellular neurosecretory cells in the rat (50), the motoneurons of the crab cardiac ganglion (123), the motoneurons of the crab stomatogastric ganglion (164), the main sensory nucleus of the trigeminal nerve (152) and a subset of respiratory neurons in the PBC (116, 119) in which rhythmic firing is abolished by a blocker of I_{CAN} (flufenamic acid) but not by a blocker of I_{NaP} (riluzole) (37, 119). Similar results were reported from investigations in dopaminergic neurons of mice substantia nigra pars compacta (102).

Different types of potassium currents also contribute to bursting activities mostly by affecting the plateaus duration and the bursting frequency. A Na⁺activated potassium current $(I_{K(Na)})$ contributes to the generation of the afterhyperpolarization potential and the maintenance of rhythmic bursts in rat neocortical neurons (48). It was also suggested that Ca²⁺-dependent potassium currents ($I_{K(Ca)}$) and a muscarine-inhibited M current (I_M) mediated by activation of the K_V7 channels may modulate the firing frequency and contribute to the burst termination in lamprey's spinal cord neurons and rat CA1 pyramidal cells (44, 53). Blocking or opening the small conductance potassium channels (SK) with apamin or 1-ethyl-2-benzimidazolinone (1-EBIO) respectively, resulted in modulation of the bursting activity in the dopaminergic neurons of the substantia nigra of rats (71).

Finally other currents, carried by mixed ionic charges, are sometimes at play and contribute mostly to the regenerative feature of this firing mode. As an example, the electrogenic sodium/potassium pump (Na^+/K^+ -ATPase) may hyperpolarize neurons and lead to activation of I_h , a cationic current activated by hyperpolarization which may provide the depolarising rebound required to reactivate the driving forces for the next burst of action potentials (32). The I_h current is often revealed by the occurrence of a depolarizing sag in response to increasing hyperpolarization. This current, carried by sodium and potassium ions has been associated to pacemaker properties in a number of neurons (for examples and review see (27, 90, 126)).

2.3 Sequence of events leading to bursts

How these various conductances interact to promote rhythmogenesis may differ from one area of the brain to another, or even within one brain structure, from neuron to neuron. Even at the single cell level, subcellular localisation of different ionic channels in different cell compartments may lead to plural bursts driving modes, as it has been proposed in Purkinje cells that showed two kinds of burst firings with distinctive waveforms; a somatically generated burst firing driven by activation of sodium persistent channels present on the soma and a dendritically generated burst firing driven by activation of LVC channels that could be found on the dendrites (47).

However, the generation of bursts presents a common sequential scheme which first requires a depolarizing current to produce the plateau potential, second, the intervention of a repolarizing process to terminate the burst and, third the need for a depolarizing

rebound to reactivate the driving force for the next burst (see Figure 1A).

Many models have been proposed to summarize the interplay of conductances leading to the generation of a bursting activity in neurons and this interplay is merely dependant on the set of channels that are comprised in the cell membrane. In masticatory, respiratory and locomotor CPGs, the observation that the depolarization prior to bursting activity occurs at membrane potential close to the potential of activation of I_{NaP} , and the fact that bursts are abolished upon application of riluzole prompted I_{NaP} as the primary driver of the neuronal bursts (see Figure $1B_1$) (20, 36, 149, 152). Since I_{NaP} is voltage-dependent, the depolarisation brought up by its activation would eventually cause its inactivation. In these cells, repolarisation and burst termination can be triggered by activation of $I_{K(Na)}$ and/or of $I_{K(Ca)}$ which contribute to the burst termination in the rat main sensory trigeminal nucleus (20) as they do in the lamprey spinal cord (44). Voltage-gated potassium channels are also present in the lamprey spinal cord and their blockade induces an increase in burst frequency (61), making them good candidates for the modulation of the burst cessation. Activation of all these potassium channels would lead to membrane hyperpolarisation which in turn would activate I_h in cells carrying this channel, thus providing for a regenerative drive. In ferret's thalamo-cortical neurons (90) and rat's hippocampal CA3 region (27), blockade of I_h with ZD 7288 abolished both the afterdepolarization and the progressive rebound depolarization and resulted in suppression of the oscillatory activity in these neurons.

In thalamo-cortical neurons, the sequence of events leading to bursting (Figure 1B₂) is thought to begin with a hyperpolarization which releases LVA calcium channels from their inactivated state making them available for activation upon depolarization which may result from activation of I_h by the prior hyperpolarization and which would also trigger firing through the recruitment of sodium and high voltage-activated calcium channels. The latter eventually activate calciumdependent potassium channels and voltage-gated potassium channels which repolarize the plasma membrane and hyperpolarize it to a level sufficient to activate I_h and drive a new boost of depolarization, leading to a renewed activation of calcium conductances (46). In this example, low voltage-activated calcium channels are clearly the primary driver of the events leading to bursting.

As to bursts driven by I_{CAN} (Figure 1B₃), Pace and colleagues (116) proposed two different mechanisms: the first one involves metabotropic mGluR5 receptors which would initiate a process leading to the release of

calcium from the intracellular stores through IP_3 receptors. The released calcium would then activate I_{CAN} channels. The second hypothesis involves calcium entry through NMDA receptors. Both mechanisms imply that a

synaptic input would activate these channels. No indication was given as to the burst termination however like in the lamprey spinal cord, potassium conductances may play that role (44, 61).

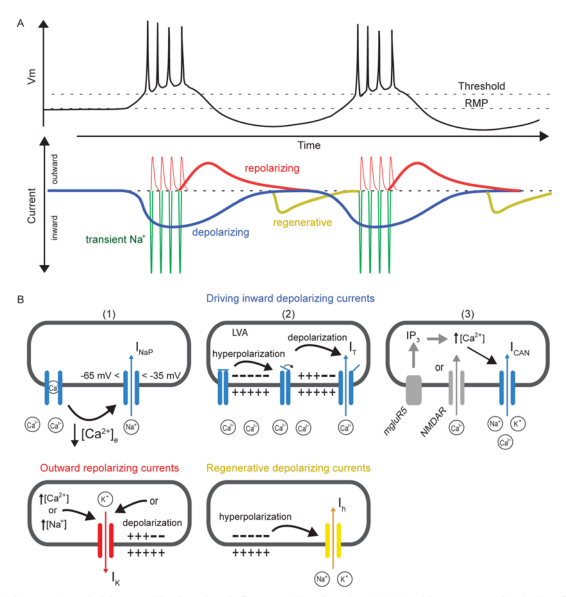


Figure 1. Ionic currents underlying repetitive bursting. A, Representation of two bursts (top) and the sequence of activation of the underlying ionic currents (bottom). Repetitive bursting generally relies on activation of a conductance that produces a sustained depolarization or plateau (blue) and "drives" the membrane potential from the resting level (RMP) to the threshold for activation of transient sodium currents (dark green) responsible for action potentials. Voltage-gated potassium channels (red) repolarize the membrane after each of these action potentials while other potassium channels activated by Ca²⁺ and Na⁺ entering the cell during depolarization and firing slowly repolarize the membrane often to a level below RMP. In many instances, the hyperpolarization triggers activation of Ih which enables regeneration of the cycle by depolarizing the membrane potential and allowing for reactivation of the driving current. **B**, Top: Different channels responsible for driving the depolarizing plateaus supporting bursting and the conditions required for their activation. Bottom: Channels required for repolarization and burst termination (red; left) and for regenerating the depolarizing drive (yellow: right).

III- ASTROCYTES ARE INVOLVED IN IONS HOMEOSTASIS AND RHYTHMOGENESIS

The most studied ions with regard to neuronal excitability and which concentrations are under strict regulation in the brain are potassium, calcium, sodium and chloride. Potassium concentration is higher in the neuronal intracellular compartment than in the extracellular space (130 mM versus 2.7-3.5 mM, respectively) (76) while the opposite is true for sodium (extracellular and intracellular concentrations are around 145 and 8-15 mM, respectively) (129). Calcium shows the highest difference with an intracellular 'resting' concentration of about 100 nM, and an extracellular concentration of 1-2 mM (35, 59, 109, 145). The extracellular chloride concentration ([Cl]_e) is around 120 mM (77) while chloride intracellular concentration is between 5-20 mM and is maintained by several molecular elements, including the cation-chloride cotransporters (NKCCs for example) and CI-HCO₃ exchangers (49, 118).

At rest or relatively low activity levels, neuronal ion pumps and transporters counteract any major ions movement and contribute to maintain the cell membrane at a relatively stable potential as well as the ionic gradient concentrations between the intra and extra-cellular compartments. However with higher activity levels, ionic concentration changes occur in both the intra- and extracellular compartments and additional mechanisms may be required to restore homeostasis. Evidence suggests that astrocytes may play a major role in controlling the extracellular concentration of some ions.

3.1 Potassium

3.1.1 Astrocytes and potassium homeostasis

During neuronal discharge, activation of the voltagegated potassium channels causes potassium ions to flow towards the extracellular space. The extracellular potassium concentration ([K⁺]_e) can increase up to a ceiling level of around 12 mM (40) in physiological conditions. Astrocytes are responsible for bringing [K⁺]_e back to resting values after an activity-driven increase. In 1966, Orkand et al. (115) introduced the concept of spatial buffering as a possible way of balancing $[K^{\dagger}]_{e}$. They proposed that the potassium accumulated during neuronal activity is taken up by neighboring astrocytes and redistributed through the syncytium to less concentrated areas. This is done mainly through the activation of gap junctions (136, 161). Indeed, it has been shown that gap junctional coupling between astrocytes is increased by elevations in $[K^{\dagger}]_e$ (34, 45), providing an anatomical support to this theory. Glial cells express

several voltage-dependent potassium channels (inward rectifier, Kir; delayed rectifier, K_d; transient A-type, K_A; Ca^{2+} -activated, K_{Ca}) (for review see references (11, 25)). However, the potassium channel that is predominantly responsible for the high potassium permeability and the maintenance of glial resting membrane potential (RMP) close to the equilibrium potential of potassium (E_K) is Kir, mostly the Kir4.1 channel. Because of the high permeability of glial membrane to potassium, this regulation of $[K^{\dagger}]_{e}$ is done mainly by a passive mechanism. However, in some cases, especially when there is an entry of sodium in the glia, an active process that relies on the action of glial Na⁺/K⁺-ATPase may be at play. In fact, according to D'Ambrosio et al. (31), both, Kir channels and the Na⁺/K⁺ pump, participate in the regulation of $[K^{\dagger}]_{e}$. Their findings suggest that neuronal and glial Na⁺/K⁺ pumps may be involved in the setting of baseline [K⁺]_e levels and its recovery rate during sustained high-frequency firing. Glial Kir channels, would also be involved in the regulation of baseline $[K^{\dagger}]_{\epsilon}$ but might not affect the rate of potassium clearance during neuronal firing. Closely linked to the Kir4.1 channels are the aquaporins-4 (AQP4): membranous channels that allow water flux through the membrane of astrocytes and that are responsible for the volume changes observed in astrocytes as well as the shrinkage of the extracellular space associated with intense neuronal activity (105). Finally, another player that helps to actively counteract the increase of [K⁺]_e is the Na⁺/K⁺/2Cl⁻ co-transporter (60, 148).

3.1.2 Extracellular potassium fluctuates with burst firing Potassium contributes to the generation of rhythmic oscillations. Indeed, using neonatal rat spinal cord in vitro preparation, Bracci et al. (18) showed that artificially rising [K⁺]_e, from a control value of 4.5 mM to over a threshold of about 8 mM, leads to sustained activation of the spinal locomotor network similar to the locomotor rhythm typically induced by serotonine and NMDA. Interestingly, they found that the suprathreshold concentration range required to elicit such sustained rhythmic motor pattern was very narrow (~ 1 mM) and when rhythmic activity was elicited with further increases (>2 mM over threshold) it was present only transiently before switching to tonic firing, then into block. Another interesting point is the facilitative effect they reported of co-applying subthreshold concentration of NMDA and serotonine that allow occurrence of rhythmic motor pattern with subthreshold (6 mM) $[K^{\dagger}]_{e}$. Similar results were found in a study carried out by Jensen and Yaari (70) on rat hippocampal slices.

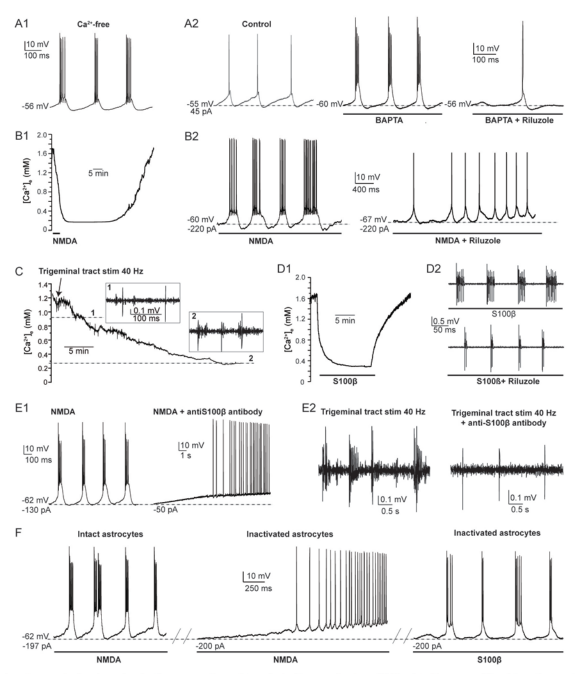


Figure 2. Involvement of astrocytes in the generation of bursts in the masticatory CPG. A1, Neurons of the trigeminal main sensory nucleus (NVsnpr) display a bursting pattern in calcium-free medium. A2, In control conditions, NVsnpr neurons fire tonically (left). Bath application of BAPTA causes one such neuron to burst (middle), and I_{NaP} channel blocker riluzole prevents the BAPTA-induced bursting in this neuron (right). B1, Local application of NMDA leads to a significant decrease in extracellular calcium. B2, NMDA-induced bursting in a NVsnpr neuron recorded in whole-cell configuration (left) is blocked by bath application of Riluzole (right). C, Electrical stimulation of the sensory trigeminal tract at 40 Hz also induces calcium depletion in the extracellular medium and bursts as seen in an extracellular recording of a NVsnpr neuron. Inset 1 shows a tonic discharge before calcium drops. Inset 2 shows a burst activity when extracellular calcium level is below 0.3 mM. D1, The astrocytic protein S100β causes extracellular calcium to diminish. D2, A NVsnpr neuron discharges rhythmically in presence of S100β (top) and S100β-induced bursts are blocked by Riluzole (bottom). E1, In whole-cell recordings, NMDA causes bursting (left) but not in presence of an antibody specifically directed against S100β (right). E2, In extracellular recordings, electrical stimulation of the sensory trigeminal tract at 40 Hz elicits bursting in control conditions (left) but not in presence of an anti-S100β antibody (right). F, In whole-cell recordings, NMDA induced rhythmic bursting under control conditions (left), but only tonic firing after inactivating a neighboring astrocyte with intracellular dialysis of BAPTA (middle). Bursting was restored with an external application of S100β (right).

The authors reported occurrence of periodic bursting in the CA3 region that spread to the CA1 when the slices were exposed to a saline with elevated potassium (7.5 mM). In a more physiological context, Marchetti et al. (96) reported that $[K^{\dagger}]_{e}$, as estimated with ion-sensitive electrodes recordings, increases to up to 8 mM and 6 mM in the spinal cord of rats during fictive locomotion electrical stimulation and by respectively. In this study, the authors did not indicate if the increases in $[K^{\dagger}]_{e}$ occurred before or after the appearance of fictive locomotion, but in a more recent study, Brocard et al. (19) using ion-sensitive electrode recordings in neonate rat spinal cord, showed that potassium increases from a resting value of 4 mM, to about 5 mM prior to the onset of locomotion-like activity induced pharmacologically (with application of NMA and serotonin) or with electrical stimulation of afferent sensory inputs and further to over 6 mM as the locomotor rhythm progresses. These evidence suggest that, at least, the early changes in [K⁺]_e can help the emergence of the rhythmic pattern since they appear prior to any detected rhythmic activity from the ventral roots, whereas the later [K⁺]_e increases could be partly consequential to the ongoing neuronal activity since they seem to be related to the increase in burst amplitude. Three of the studies mentioned above were performed on the same preparation (spinal cord, in vitro), in the study of Bracci et al., $[K^{\dagger}]_{e}$ was artificially controlled whereas in the others two, the values of $[K^{\dagger}]_{e}$ reported were obtained in more physiological conditions. However, the three studies reported similar values of [K⁺]_e correlated to occurrence of rhythmic activity. Extracellular potassium increases were also observed in the lamprey spinal cord during fictive swimming (158). The authors reported a two-components $[K^{\dagger}]_e$ increase consisting in phasic increases of $[K^{\dagger}]_e$ of around 0.2 mM linked to the ventral root discharges on the ipsilateral side surimposed on a slow elevation (varying between 0.08-0.40 mM) of the baseline level of $[K^{\dagger}]_e$ that parallels the initiation of ventral root burst activity. The authors did not clearly state if the onset of the slow elevation precedes that of the detected rhythmic activity, but their observation that the general form of the $[K^{\dagger}]_{e}$ -curve closely resembles that of the bursts on the ipsilateral side suggests that the phasic $[K^{\dagger}]_e$ increases may be a consequence of neuronal activity. Similar results have been reported in the cat ventrolateral medulla in parallel with central respiratory activity in the phrenic nerve (125). In this case, $[K^{\dagger}]_{e}$ started to rise prior to the discharge of action potentials, thus the authors proposed that the efflux of potassium was produced as a consequence of synaptic transmission. Finally, local transients (between 1-2 mM) in $[K^{\dagger}]_{e}$ were also seen

during cortical slow sleep oscillations and spike-wave seizures (4).

Extracellular potassium likely interacts with the conductances that shape the bursts. Indeed, low concentrations of TEA or increased $[K^+]_e$ have been shown to prolong burst duration and to increase oscillations amplitude in the supraoptic nucleus of rats (86) probably by preventing burst termination. A computational model of Hb9 cells suggests that increase in $[K^+]_e$ does not upregulate I_{NaP} in locomotor CPG circuits (19). However, recordings from CA1 pyramidal neurons in hippocampal slices revealed a TTX-sensitive persistent inward current in these neurons that was reversibly enhanced when $[K^+]_e$ was raised (141). It has also been shown that elevated $[K^+]_e$ increases the conductance of the channels mediating I_h (143).

3.1.3 Evidence for spatial buffering of potassium during bursting

Is there any evidence that the astrocyte-driven spatial buffering of potassium takes place along with rhythmic firing in physiological conditions? Working on brainstem slice preparations of neonatal rats, Schnell et al. (137) reported periodic fluctuations of the membrane current of astrocytes occurring in phase with the rhythmic discharges of PBC neurons. Blockade of Kir channels with barium (Ba²⁺) decreased the amplitude of the periodic membrane current fluctuations by more than 50% suggesting that these currents partially reflect the periodic uptake of the elevated extracellular potassium around the astrocytes with each fired burst. Such potassium buffering seems to occur along with largescale oscillations as well. In the cerebral cortex of anesthetized cats, a periodic increase in intraglial K⁺ concentrations ([K⁺]_i) occurs simultaneously to the depolarizing phase of the slow oscillations (3).

Impairment of spatial buffering may be involved in pathologies like epilepsy. Epilepsy is characterized by periodic seizures resulting from abnormal neuronal hyperexcitability and high synchronicity. Epileptiform activity is generally initiated in restricted areas of the brain but can spread to other parts (8, 106, 121, 151). It has been known for more than 4 decades that extracellular potassium increases during epileptogenesis (99), and many evidence suggest a role for astrocytes in [K⁺]_e regulation in this pathology. For instance, a study by Kivi et al. (75) suggests an impairment of potassium buffering in hippocampal CA1 slices from epileptic rats. Indeed, an application of Ba²⁺ did not affect the level of extracellular potassium in epileptic slices with Ammon' Horn sclerosis. In sclerotic tissues, astrocytes show lower densities of Kir channels (62). Beside Kir channels, many investigations also suggest that astrocyte-expressed aquaporins may play a significant part as they are responsible for the water flow, therefore volume changes in the extracellular space and ultimately the $[K^{\dagger}]_{e}$. Indeed, epilepsy has been associated to abnormal expression of aquaporin AQP4 in mice hippocampus (85) and human cortex (97).

3.2 Calcium

3.2.1 Astrocytes and calcium homeostasis

Neuronal activity is associated to a decrease of the extracellular concentration of calcium [Ca²⁺]_e that occurs in parallel to the rise in [Ca²⁺], due to calcium influx into the cells through voltage- and ligand-gated ion channels. Contrarily to potassium, much less is known about the role of glia in the regulation of [Ca²⁺]_e. It is however known that in many brain areas, astrocytes possess voltage-gated calcium channels (25, 82) and express calcium sensors (24). Furthermore, direct evidence show that astrocytes can effectively sense the level of calcium in the extracellular compartment. Indeed, Zanotti and Charles (163) have shown that astrocytes exposed to low [Ca²⁺]_e respond with increases in [Ca²⁺]_i that propagate intercellularly as Ca²⁺ waves. This is consistent with the observation that decreased [Ca²⁺]_e promotes the opening of hemichannels that mediate electrical coupling of neurons and glial cells (150). This low calcium-mediated intracellular calcium rise, however, relies on a release of calcium from intracellular stores since it is inhibited by thapsigargin (163). There is evidence that the astrocytic Ca²⁺ waves modulate the firing frequency of ganglion cells in dissected eyecup retinas neurons (108). Lian and Stringer (87) have shown that during induced spreading depression in the rat cortex, the [Ca²⁺]_e remained low for a longer time-lapse than normal and the shape of the recovery curve was altered when astrocytes activity was artificially inhibited with fluorocitrate (FC) and fluoroacetate (FA). Indeed, under control conditions, the [Ca²⁺]_e recovery curve presents a phase of transient overshoot indicative of an active extrusion of calcium from cells in an amount greater than that which had moved into the cells. The overshoot phase was lacking with the use of FC/FA suggesting that the failure of astrocytes to actively extrude calcium may account for the longer time required to restore the level of extracellular calcium.

3.2.2 Extracellular calcium fluctuates with burst firing The extracellular concentration of calcium also seems to be determinant for bursting in a number of brain areas. Although, increases in $[Ca^{2+}]_e$ can lead to bursting as shown by Formenti *et al.* (46) in rat thalamic neurons, bursting is most often associated to decreases of $[Ca^{2+}]_e$. *In vivo*, bursting in the neocortex during seizures is

associated to decreases in [Ca²⁺]_e that occur prior to the onset of phasic activity (59). In vitro, artificial reduction of [Ca²⁺]_e have been associated to neuronal bursting in preparations of the rat supraoptic nucleus (86), the hippocampus (146), medullary respiratory neurons (36. 73, 114), spinal neurons (19), and neurons of the trigeminal main sensory nucleus (Figure 2A₁) (20, 101, 152). In most of these studies, the medium used was calcium free. In the trigeminal main sensory nucleus, even locally restricted changes induced by extracellular application of the Ca²⁺ chelator BAPTA convert tonic firing into rhythmic bursting (Figure 2A₂). More interestingly, stimuli like NMDA (locally applied) or stimulation of sensory inputs to the nucleus, which efficiently elicit bursting in these neurons (Figure 2B2, 2C and 2E₂) are also associated to a decrease in [Ca²⁺]_e (close to 0.89 and 1.1 mM, respectively) (Figure $2B_1$ -C). In the spinal cord, Brocard et al. (19) reported 1.03 mM as the extracellular Ca2+ concentration below which they observed locomotor-like activity when they electrically stimulated the ventral funiculus, whereas in the main sensory trigeminal nucleus the [Ca²⁺]_e needed to drop beneath 0.4 mM for rhythmic activity to appear (101). These differences may reflect differences in age, structures, or electrodes placement between these two studies. In neurons of the trigeminal main sensory nucleus, bursting elicited by NMDA or BAPTA applications is driven by I_{NaP} (Figure $2A_2$ and $2B_2$). Depletion of extracellular calcium shifts the I-V curve of I_{NaP} towards more hyperpolarized potentials (86, 101, 149) and from predictions based on a computational model of Hb9 cells, it appears that a slight shift of I_{NaP} activation suffices to switch the neurons firing pattern from tonic to rhythmic bursting (19). The work by Armstrong (5) in the squid giant axon suggests that Ca2+ may favor closing of the channel by occupying its pore. In its absence, closing slows considerably or does not occur. This effect of calcium may also help understand the voltage dependency of the activation range of $I_{\rm NaP}$ since the ability of calcium to enter the pore, and block it, is reduced as voltage is driven negative.

Unlike sodium persistent channels, activation of LVA Ca²⁺ channels is promoted with rises in [Ca²⁺]_e. This effect may result from the fact that these channels are activated at more hyperpolarized potentials. Increases of [Ca²⁺]_e hyperpolarize thalamic neurons and shift the conductance–voltage relationship of their LVA Ca²⁺ channels to the right (46). The hyperpolarization, which may result from a masking of the negative fixed charges on the membrane surface by the calcium ions facilitates activation of the LVA channels, whereas the increased Ca²⁺ driving force favors appearance of low threshold Ca²⁺ spikes and eventually bursting.

3.2.3 Astrocyte-mediated extracellular calcium depletions during bursting

Although often postulated, direct evidence of astrocytic regulation of extracellular calcium were only recently obtained. We recently showed that an astrocytic protein, S100 β , reduces $[Ca^{2+}]_{\epsilon}$ (Figure 2D₁) and induces I_{NaP} driven bursting in neurons of the trigeminal main sensory nucleus (Figure 2D₂) (101). Protein S100β was proven to be released from astrocytes (155) after elevation of cytosolic Ca²⁺ (33), but the exact mechanisms underlying its release or secretion have not been deciphered yet. Some studies (26, 134) suggested that these mechanisms may involve the activation of glutamate mGluR3 and/or A1 adenosine receptors, while others (134) have ruled out involvement of Cx43 hemichannels. S100ß exerts many actions, one of which is calcium chelation (94, 95). In support of this role, we have observed that blockade of endogenous \$100\beta with application of an antibody directed against it prevents bursting and Ca²⁺ decreases induced in the trigeminal sensory nucleus with NMDA or sensory fibers stimulation (Figure 2E₁₋₂ and (101)). These findings strongly suggest a prominent role for astrocytes in rhythmogenesis. This is further supported by the finding that preventing activation of astrocytes with intracellular dialysis of BAPTA in one of several connected astrocytes led to cessation of NMDA-induced bursting (Figure 2F and (101)) in adjacent neurons. Bursting blocked by this procedure could be restored by exogenous application of \$100\beta (Figure 2F right; and (101)). Thus, we propose that in the trigeminal main sensory nucleus, which is postulated to form part of the core of the masticatory CPG, rhythmic activity results from activation of astrocytes by incoming sensory glutamatergic inputs (101) leading to release of S100B which in turn binds extracellular calcium, causing a decrease in $[Ca^{2+}]_e$ and a consequent shift in I_{NaP} activation curve which facilitates bursting (Figure 4). This study constitutes the first demonstration that directly link astrocytic S100β to neuronal firing pattern in individual cells (101). However, other evidence suggest that S100β can also play a role in large-scale oscillations. In two studies using S100\beta knock-out mice, no significant differences were found in the spontaneous oscillations detected in the hippocampus and the neocortex as compared to wildtype mice (133, 134). However, these oscillations diverged during kainate-induced seizures. Under these conditions, there was a significant increase in the amplitude of gamma waves (133, 134) that was correlated with increased level of extracellular S100ß in the wildtype (134), suggesting that S100β may contribute to gamma oscillations in the hippocampus. Gamma oscillations are high frequency waves (30-80Hz) associated with attentive behavioural states that can be found in the neocortex and the hippocampus. Evidence of S100 β implication in these rhythmic events support our data showing an important role for S100 β in rhythmogenesis in the masticatory GPG.

Several other findings link changes in astrocytic functions to dysregulation of extracellular calcium and brain pathologies. In kainate-induced epilepsy in rats for instance, immunocytochemical investigations have shown an upregulation of L-type voltage-gated calcium channels α2 subunit in astrocytes surrounding the lesion (160). No significant changes were reported in neurons. In the hippocampi of patients who suffered from temporal lobe epilepsy associated with Ammon's horn sclerosis, astrocytes were also strongly immunoreactive for $\alpha 1c$ subunits (42). Some of the voltage-gated calcium channels have a relatively high threshold of activation and given the very negative membrane potential of healthy astrocytes it is unlikely that they can be depolarised enough to activate these channels. Nevertheless, it has been shown using brain slices or cultures that astrocytes from human sclerotic epileptogenic hippocampal seizure foci display more depolarized membrane potential (around 255 mV) comparatively to astrocytes from nonsclerotic tissue (around -75 to -80 mV) (112) and express higher TTXsensitive Na⁺ channel density allowing them to generate potential-like responses with membrane depolarization (17, 112). In these conditions, one should expect such upregulations of L-type voltage-gated calcium channels subunits to lead to an increased flow of calcium into astrocytes, potentially causing a decrease in [Ca²⁺]_e. Parkinson's disease may be another example of pathology that results from an inefficient regulation of calcium levels in the extracellular space. Symptoms of Parkinson's disease include neuronal death and excessive synchronicity of beta waves in the cortex and in the basal ganglia (22, 56, 153). Glial cells play a significant role in the ontogenesis of that disease (55). In a mouse model of (1-Methyl-4-Phenyl-1,2,3,6-TetrahydroPyridine)induced Parkinson's disease, the number of S100βpositive astrocytes increases in the striatum and the substantia nigra immediately after treatment with MPTP, but declines after seven days (104). Increases in the levels of S100ß were also reported in midbrain slices and cerebrospinal fluid of patients who suffered from Parkinson's disease (135). S100β is a calcium-binding protein (94, 95) and it is possible that its transitory accumulation in the extracellular space leads to a chelation of calcium. A precursor sign of the Parkinson's disease could then be a transient decrease of calcium in the extracellular space.

3.3 Sodium

3.3.1 Astrocytes and sodium homeostasis

As neurons fire, there is a decrease in the extracellular sodium concentration ([Na⁺]_e) due to influx of sodium ions into the cells through voltage- and ligand-gated ion channels (41). Indeed, in a study carried out on hippocampal slices, Karus et al. (74) reported that during bursts, the neuronal intracellular concentration of sodium ([Na⁺]_i) increases to up to 22 mM from a resting baseline value of about 14 mM and recovers to around baseline level between bursts. Intra-neuronal concentrations of sodium need to be tightly controlled in order to maintain cells integrity and excitability and also because extrusion of sodium ions by the Na⁺/K⁺-ATPase is an energy-expensive process. How do astrocytes participate in sodium homeostasis? Astrocytes express voltage-gated Na⁺ channels (for a review about sodium homeostasis and signaling in astrocytes see reference (128)) which allow for a small but steady influx of Na⁺ ions to ensure the maintenance of cytoplasmic Na⁺ at concentrations required for proper functioning of the glial Na⁺/K⁺-pump (142). Moreover, sodium signalling concomitant with neuronal activity has been shown to occur in astrocytes. Indeed, combining somatic wholecell patch-clamp recordings with quantitative sodium imaging with the sodium-sensitive fluorescent indicator dye (sodium-binding benzofuran isophthalate), Bennay et al. (15) showed that short bursts of synaptic activity resulted in glial sodium signals of up to 9 mM in cellular branches of cerebellar Bergmann glial cells. These sodium increases persisted for tens of seconds. Sodium may also build up in astrocytes following excitatory neuronal activity because of the glial glutamate transporter which carries 3 Na⁺ ions with each glutamate molecule (80). In cultures of cortical astrocytes, uptake of glutamate produces intercellular sodium waves that occur in parallel with calcium waves following stimulation of a single astrocyte. Sodium underlying these waves has to come from the extracellular space since it could not be released from the intracellular stores as is calcium. Thus, at sites of intense activity, the consequent lowering of extracellular sodium could weaken the drive for this ion towards the neuronal intracellular compartment. Besides the glutamate transporter, the glial Na⁺/K⁺-pump, which is involved in $[K^{\dagger}]_{e}$ homeostasis, can also greatly impact on neuronal sodium homeostasis. Indeed, blocking astrocytes metabolism during basal activity using sodium-fluoroacetate results in an increase of baseline intracellular sodium of around 4 mM in hippocampal neurons and by around 12 mM in astrocytes (74). The authors attribute the larger increase in astrocytes to the weakening of the glial Na⁺/K⁺-ATPase. Under conditions of increased network activity, blocking of astrocytes metabolism increases even more the baseline intracellular sodium in both cells types but also produces a nearly fivefold prolongation of individual epileptiform bursts and similar increase in the number of population spikes per burst (74). The authors proposed that the increased neuronal excitability results, in part, from reduced glutamate uptake due to reduced activity of the glutamate transporter that greatly depends on the transmembrane sodium gradient in astrocytes which in turn depends on the activity of the Na⁺/K⁺-ATPase. Furthermore, since the glial Na⁺/K⁺-ATPase is also involved in the clearance of extracellular potassium following intense neuronal activity, its weakening reduces potassium intake thus increasing neuronal excitability. No measurement of extracellular sodium have been made in parallel in this study, however, these evidence of astrocytes involvement in the control of neuronal [Na⁺]_i suggest that even indirectly, by achieving other basic cellular functions, astrocytes may participate in the homeostasis of extracellular sodium.

3.3.2 Extracellular sodium and bursting

Studies addressing the direct role of sodium ions in bursting are relatively scarce. Much of the studies addressed the involvement of sodium channels rather than the role of the ionic sodium gradient per se. One study by Li and Hatton (86) carried out on the magnocellular neurons of the hypothalamic supraoptic nucleus (SON), showed that both, $[Na^{\dagger}]_{e}$ and $[Na^{\dagger}]_{i}$ are determinant for the low calcium-mediated burst firing observed in a subpopulation of these neurons. They examined the effect of reducing the extracellular sodium and reported that this treatment abolished or reduced bursting in all tested cells. Finally, even though no study has formally measured the changes in extracellular sodium during rhythmic activities, [Na⁺]_e is likely to decrease because of all the voltage and ligand-activated channels allowing movements of Na⁺ ions from the space towards the extracellular intracellular compartment. However, the question remains as to the extent of sodium depletion and how such decreases may influence other conductances involved in bursting.

3.3.3 Evidence for synchronized sodium signaling in astrocytes somata during bursting

In the study cited above, Karus *et al.*, (74) showed that during neuronal recurrent bursting, astrocytes showed an increase of about 2.9 mM in [Na⁺]_i that occurred in parallel with the transient [Na⁺]_i increase seen in neurons. In about 30% of the astrocytes, this increase was followed by an undershoot below baseline. The increases of astrocytic [Na⁺]_i observed during neuronal bursts were accompanied by recurring membrane

depolarisations of about 10 mV and were paralleled by recurring transient increases of $[K^{\dagger}]_e$ of up to 2 mM. Between bursts, the astrocytic $[Na^{\dagger}]_i$ recovered to resting baseline values. Using several pharmacological tools, the authors concluded that sodium signals in astrocytes result from dual contribution of two opposing mechanisms: an influx of sodium ions caused by the sodium-dependent glutamate uptake and an efflux of sodium ions brought out by the elevated $[K^{\dagger}]_e$ —induced activation of the Na^{\dagger}/K^{\dagger} -ATPase.

3.4 Chloride

The movements of chloride ions in response to neuronal activation are more complex than the other ions (41). Probably, since it is the main permeable anion, chloride ions movements are partly determined by balancing measures in relation to fluctuations of the cations gradient concentration. Dietzel *et al.* (41) reported that during stimulus-induced self-sustained afterdischarges (SAD) in neurons of the sensorimotor cortex of cats, [Cl⁻]_e always showed an average increase of 7 mM (when measured at a depth of 1 mm) often preceded by an initial small decrease. They observed that the maximum increase appears to coincide with the end of the ictal period.

3.4.1 Astrocytes and chloride homeostasis

Chloride channels exist in astrocytes and include volumesensitive chloride channels (157). The latter participate in ions homeostasis mainly through the process of astrocyte swelling (30). By swelling, astrocytes change their volume and thereby control the concentration of several ions in both the intra and extracellular space. In addition to potassium-mediated machinery which intervenes in astrocyte swelling (39, 107), an inward rectifier chloride current (14, 81) and Na⁺/K⁺/Cl⁻ cotransporter are also activated (69). The participation of chloride channels to astrocytic volume change might result from their association with actin proteins which form the cytoskeleton (157). Bikson and collaborators (16) tested the effect of DNDS (4,4'-dinitrostilbene-2,2'-disulfonic acid), a glial Cl channel blocker (103), on the duration of low calcium-mediated bursts in rat pyramidal CA1 cells and found that high concentration of DNDS caused a 6-to 10-fold increase in burst duration suggesting that glial uptake of chloride ions contributes to the burst termination.

3.4.2 Extracellular chloride and bursting

Working on the leech's Retzius neurons, Beck and colleagues found that decreasing [Cl]e (1 mM and less) induces a sustained membrane depolarization and recurrent bursting activity that were accompanied by recurrent rises in intracellular Ca²⁺ which oscillated in synchrony with the bursts. Using a chloride-free and calcium-free medium, they determined that the intracellular calcium rises were presumably due to Ca2+ influx through voltage-dependent Ca2+ channels, since they could no longer be seen in this medium. Moreover, the bursting activity was not affected by depletion of intracellular calcium but was inhibited by saxitoxin, a sodium channel blocker, suggesting that it relies on I_{NaP} (13). In rat hippocampal slices, spontaneous (63) or stimulus-evoked (9) bursting discharges also developed when the [Cl]e was decreased. These cellular discharges were accompanied for a time with synchronous oscillations in the field potential. However, longer exposure to low chloride medium, desynchronized the firing activity of neuronal populations in the CA1 region of hippocampal slices which may explain the antiepileptic effect of chloride-cotransport blockade by furosemide (64). Reciprocally, it has been shown that calcium-free mediated bursting occurs concomitantly with large decreases in [Cl]_e (58).

3.5 Synergistic interactions and rhythmogenesis

Changes in the ionic gradient concentration of the main ions that regulate brain activity exert tremendous effects on neuronal excitability and are determinant of the neuronal firing mode. By their concerted actions on the rhythmogenic conductances, they may act in synergy to promote rhythmogenesis. Indeed, Brocard *et al.* (19) reported that such a synergistic effect exists between the $[Ca^{2+}]_e$ reduction and the $[K^+]_e$ increase for bursts generation in the locomotor CPG (Figure 3).

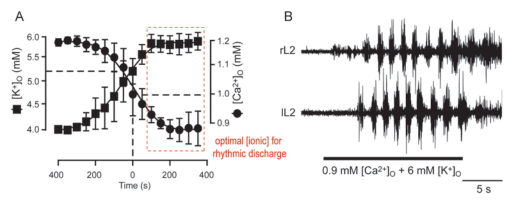


Figure 3. Example of synergistic interactions between potassium and calcium in rhythmogenesis. A, Evolution of extracellular concentration of calcium (circle) and potassium (square) before and after the onset (at 0 ms) of a locomotor-like bursting episode. Optimal concentrations of calcium and potassium for a bursting activity are emphasized by the dashed rectangle. A rhythmic activity appears at concentrations of calcium and potassium close to 0.9 and 6 mM, respectively. **B**, Extracellular recordings in the right and left L2 segments (rL2 and IL2) of rat spinal cord along with application of 0.9 mM [Ca²⁺], and 6 mM [K⁺]. Modified with permission from (19).

The authors stated that both $[Ca^{2+}]_e$ and $[K^+]_e$ concurrently changed before any rhythmic activity was detected from ventral roots suggesting that these changes are not just a consequence of the neuronal activity, but are part of the cause for the rhythmic pattern observed. They confirmed this synergistic interaction by artificially manipulating both ions concentrations while recording neuronal activity and reported that concomitant reduction of [Ca²⁺]_e to 0.9 mM and increase of $[K^{\dagger}]_e$ to 6 mM (the same values reported with physiologically-induced changes in concentration for both ions) elicited bursts in 25% of the recorded neurons. A similar interaction between both ions concentrations reported this time as a "Ca²⁺/K⁺ antagonism" is also encountered in mice rhythmogenic inspiratory active preBötzinger complex (117). The authors indeed showed that the rhythm generated in 3 mM K⁺ and 1 mM Ca²⁺ is depressed following a modest rise of Ca²⁺ and restored when K⁺ is subsequently raised. Ultimately, they proposed that the optimal window to obtain long term stable slice rhythm would be 0.75-1mM for calcium and 4-6 mM for potassium, which seems in accordance with the values obtained for either ions in more physiological conditions (4, 19, 96, 125). How do all of the elements converge to allow the emergence of an oscillatory activity? From the reports examined here, it seems that the starting point is the increased firing that causes an elevation of extracellular potassium. According to Hounsgaard and Nicholson (65), an individual burst is initiated when intense neuronal firing results in a local increase in $[K^{\dagger}]_e$ that would in turn depolarize neighboring neurons. Increased activity of these neighboring neurons would boost the $[K^{\dagger}]_e$ to a further

extent. This increased activity and [K⁺]_e would directly activate the astrocytes and cause them to start the spatial buffering of potassium to prevent local build up of extracellular K⁺ that would cause a depolarizing block and cessation of cell activity. Spread of K⁺ will be limited only to astrocytes that are coupled. Thus, gap junctional coupling between astrocytes may provide a mean to limit spreading of activity to a specific neuronal population as well as an explanation for the observed synergistic interaction between ions. As already stated, extracellular accumulation of K⁺ increases gap-junctional coupling between astrocytes. Interestingly, Scemes and Spray (136) found that this increase in dye-coupling long outlasted the exposure to elevated potassium, suggesting that the effect of potassium on coupling may be indirect. Indeed, the elevated extracellular K⁺ causes astrocytic membrane depolarization and leads to the influx of Ca²⁺ through L-type voltage-activated Ca²⁺ channels (34, 43, 147), promoting increases in [Ca²⁺]_i. Blocking the influx of calcium with the L-type calcium channel blocker nifedipine or potentiating this influx with Bay-K-8644, prevented and potentiated the K⁺-induced increase in coupling, respectively (34). At last, they provide evidence that the K⁺-induced increased coupling relies on the activation of a CaMkinase by the intracellular calcium rise since it can be prevented by the calmodulin antagonist calmidazolium and by the inhibitor of CaMkinases, KN-93. The authors hypothesized that the increased coupling might result from an increased number of active channels within gap junction plaques. However, only very large increases of extracellular potassium (50 mM), much larger than those observed even with intense neuronal activity 8-12 mM (140)

yielded an elevation of [Ca²⁺]_i in astrocytes, suggesting that voltage-gated Ca²⁺ influx would be unique to pathological conditions associated with deregulations of [K[†]]_e. However, the accumulation of sodium in astrocytes during intense excitatory discharge may also cause a calcium entry by reversing the function of the sodium/calcium exchanger (52, 124, 127) and consequently lead to an extracellular decrease of Ca²⁺. Other mechanisms may also account for the decreased calcium level in the extracellular compartment that occurs in conjunction with rhythmic firing. Astrocytes have a wide array of ionotropic and metabotropic receptors (25), the activation of which leads to intracellular rises of calcium and glutamate-induced calcium waves in astrocytes have been commonly reported (1). As described above, data from our laboratory suggest that glutamatergic activation of astrocytes may also cause release of the protein S100ß from astrocytes, leading to extracellular calcium depletion, subsequent I_{NaP} activation and burst firing (101). Figure 4 summarizes how astrocytes may participate in emergence and synchronization of rhythmic firing in NVsnpr neurons. The proposed model rests on the fact that NVsnpr astrocytes are activated by glutamatergic afferent inputs to this nucleus, but only if they reach a sufficient level of activity. At high activity level, glutamate or K⁺ released from these inputs activate astrocytes and cause release of S100\u03b3. The extent of neurons exposed to the released \$100\beta will depend on the extent of coupled astrocytes. Among these neurons, only those having I_{NaP} will change their firing. In young rats (P8-12), only 38% of the dorsal NVsnpr neurons perfused with a calcium-free medium showed bursts firing and, bursting and non-bursting cells displayed distinct morphological characteristics and significantly different input resistance and membrane capacitance (152). However, using older rats (P16), Brocard et al. showed that 85% of dorsal NVsnpr neurons had intrinsic bursting properties in calcium-free medium and bursting relied on I_{NaP} , in all tested cells (20). We postulate that the extent of the active network would likely be determined by the population of cells (neurons and astrocytes) that share the same input, and since, the primary afferent inputs are somatotopically organised, only functionally related cells would share the same Eventually, cessation of primary afferents activation would stop the network activation, or excessive stimulation would depolarize the neurons beyond the range of I_{NaP} activation.

IV- CONCLUSION

The classic concept in which brain function relied exclusively on neuron-to-neuron communication was shattered by evidence in which glial cells switched from the role of brain glue to active information processors expressing a large spectrum of ion channels and receptors. This review recounts the progress made in understanding the prominent role that glial cells play in the generation and maintenance of rhythmic oscillations. Glial cells have proven to be essential partners in coordinating the numerous elements participating to rhythmogenesis through a cross-talk with surrounding neurons. The control of extracellular concentration of ions necessary for rhythmic activities to occur clearly depicts that dialogue.

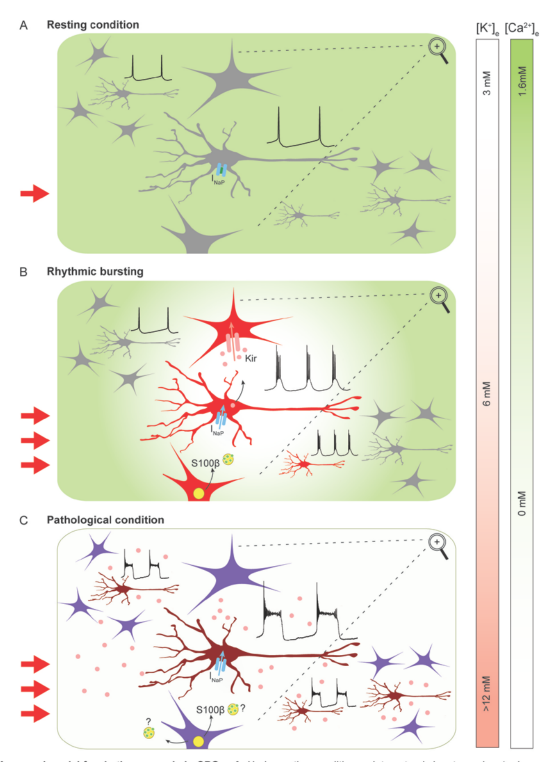


Figure 4. A general model for rhythmogenesis in CPGs. A, Under resting conditions, glutamatergic inputs are low (red arrow), astrocytes (grey) are at resting activity levels, $[Ca^{2*}]_e$ is elevated and $[K^*]_e$ is low. The I_{NaP} channels are obstructed by calcium (green dot) and neurons fire tonically. B, As the glutamatergic inputs increase, $[K^*]_e$ (pink dots) rises, astrocytes get activated (red), release S100β (yellow) and initiate spatial buffering through Kir channels. Protein S100β chelates calcium and causes $[Ca^{2*}]_e$ to drop. As a consequence, I_{NaP} channels are freed and drive bursting in neurons (red). The bursting population is limited to the extent of coupled astrocytes forming a syncytium. C, Under pathological conditions, astrocytes (purple) fail to initiate spatial buffering in the presence of intense neuronal activity, leading to an accumulation of potassium (pink dots) and spread of excitation and depolarization block among neurons. The level of S100β in the extracellular space may increase and further lead to neuronal hyperexcitability by decreasing $[Ca^{2*}]_e$.

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