Studies of Respiratory Rhythm Generation Maintained in Organotypic Slice Cultures

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Studies of Respiratory Rhythm Generation Maintained in Organotypic Slice Cultures

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A Dissertation presented to the Graduate Faculty of the College of William and Mary in Candidacy for the Degree of Doctor of Philosophy

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The College of William and Mary
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This Dissertation is submitted in partial fulfillment of the requirements for the degree of

Doctor of Philosophy

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ABSTRACT

Breathing is an important rhythmic motor behavior whose underlying neural mechanisms can be studied in vitro. The study of breathing rhythms in vitro has depended upon reduced preparations of the brainstem that both retain respiratory-active neuronal populations and spontaneously generate respiratory-related motor output from cranial and spinal motor nerves. Brainstem-spinal cord en bloc preparations and transverse medullary slices of the brainstem have greatly improved the ability of researchers to experimentally access and thus characterize neurons important in respiratory rhythmogenesis. These existing in vitro preparations are, however, not without their limitations. For example, the window of time within which experiments may be conducted is limited to several hours. Moreover, these preparations are poorly suited for studying subcellular ion channel distributions and synaptic integration in dendrites of rhythmically active respiratory neurons because of tortuous tissue properties in slices and en bloc, which limits imaging approaches. Therefore, there is a need for an alternative experimental approach. Acute transverse slices of the medulla containing the preBötzinger complex (preBoC) have been exploited for the last 25 years as a model to study the neural basis of inspiratory rhythm generation. Here we transduce such preparations into a novel organotypic slice culture that retains bilaterally synchronized rhythmic activity for up to four weeks in vitro. Properties of this culture model of inspiratory rhythm are compared to analogous acute slice preparations and the rhythm is confirmed to be generated by neurons with similar electrophysiologic and pharmacological properties. The improved optical environment of the cultured brain tissue permits detailed quantitative calcium imaging experiments, which are subsequently used to examine the subcellular distribution of a transient potassium current, I_A, in rhythmically active preBoC neurons. I_A is found on the dendrites of these rhythmically active neurons, where it influences the electrotonic properties of dendrites and has the ability to counteract depolarizing inputs. These results suggest that excitatory input can be transiently inhibited by I_A prior to its steady-state inactivation, which would occur as temporally and spatially summating synaptic inputs cause persistent depolarization. Thus, rhythmically active neurons are equipped to appropriately integrate the activity state of the inspiratory network, inhibiting spurious inputs and yet yielding to synaptic inputs that summate, which thus coordinates the orderly recruitment of network constituents for rhythmic inspiratory bursts. In sum, the work presented here demonstrates the viability and potential usefulness of a new experimental model of respiratory rhythm generation, and further leverages its advantages to answer questions about active currents in dendrites that could not previously be addressed in the acute slice model of respiration. We argue that this new organotypic slice culture will have widespread applicability in studies of respiratory rhythm generation.
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INTRODUCTION

Our understanding of the neural mechanisms for breathing has advanced exponentially because of studies exploiting reduced *in vitro* preparations, which broaden the scope of feasible experiments by providing a tractable laboratory model of the behavior (respiration). *In vitro* breathing models retain core brain circuitry necessary for the production of respiratory-related rhythms while improving access to neuronal populations of interest for recording and (in some cases) manipulation. Respiratory rhythmogenesis has been studied *in vitro* using acute brainstem-spinal cord and medullary slice preparations for more than 30 years (Funk and Greer, 2013; Smith et al., 1991; Suzue, 1984). Despite their numerous advantages, these preparations are not amenable to some contemporary viral transfection techniques nor imaging methodologies. Acute preparations containing respiratory neural circuits are poorly suited for these methods because they remain only viable for hours, whereas viral transfection can take days. Also, acute slices are hundreds of microns thick and even the most sophisticated imaging methods are not practicable at depths exceeding 100 µm (at the most), which forecloses our ability to measure the majority of respiratory neurons (or glia) at depths that exceed 100 µm. In contrast, organotypic culture models of respiratory rhythm generation may provide an alternate rhythmically active platform suited to long-term experimentation and imaging (on the order of
days and weeks), allowing for a new array of techniques to investigate the biophysical mechanisms giving rise to the respiratory rhythm.

Throughout the 1980s, mammalian respiration was studied primarily in vivo using decerebrate cats (Eldridge et al., 1976, 1981; Foutz et al., 1989; Piafferiche et al., 1994; Ramirez et al., 1998; Richter, 1982; Schwarzacher et al., 1995; Smith et al., 1989). The study of respiratory rhythm in vivo suffered from an inability to precisely control the recording milieu, and required the use of anesthetics such as pentobarbital, which significantly enhance the strength of chloride-mediated synaptic inhibition (thus changing the network properties). Reduced in vitro preparations that preserved respiratory rhythm-generating functionality were developed using neonatal rodents, beginning with the brainstem-spinal cord en bloc preparation (Smith and Feldman, 1987; Smith et al., 1990; Suzue, 1984). In vitro preparations were further reduced through a series of serial transections that identified the location of the preBötzinger complex (preBötC)--the core rhythm-generating network necessary for inspiratory-related motor output (Smith et al., 1991). Transverse slices of the medulla containing the preBötC, which also retain respiratory related hypoglossal (XII) motoneurons and premotor neurons, produce a bilaterally synchronized rhythm that can be monitored as motor output from the hypoglossal cranial nerve rootlets that are also captured in the slice (Figure I.1). If properly sectioned (Ruankittisakul et al., 2006, 2011, 2014), the rostral surface of these slices exposes neurons in the preBötC for detailed recording experiments, and for over 25 years acute transverse slices containing the preBötC have been utilized to define the behavior, membrane properties, synaptic connectivity, and
Figure 1.1 The in vitro slice preparation of the preBötzinger complex. Top-left panel: brainstem of a newborn mouse and diagram showing the rostral and caudal locations of slice transection (dotted lines). Top-right panel: Cycle-triggered average of preBötC bilaterally synchronized rhythmic calcium activity imaged using the membrane-permeable fluorescent calcium indicator dye, Fluo-8AM. Green and red areas correspond to active regions in the slice. Bottom-left panel: a schematic diagram of the slice, showing locations of the bilateral preBötC, nucleus ambiguous (Amb), hypoglossal motor nucleus (XII) and hypoglossal cranial nerve rootlet (XIIn). Bottom-right panel: inspiratory-related motor output recorded through a suction electrode on XIIn.

The genetic identity of interneurons involved in respiratory rhythmogenesis (Bouvier et al., 2010; Feldman et al., 2013; Gray et al., 1999; Rekling et al., 1996). However, this acute slice preparation has inherent limitations. Acute slice preparations are at best useful for several hours (Funk and Greer, 2013), precluding chronic pharmacological experiments or the use molecular techniques requiring long incubation periods (e.g., viral transfection). They are also poorly suited for
assaying subcellular distributions of ionic membrane conductances or synaptic integration occurring in dendrites.

In other regions of the brain, the integrative properties of dendrites have been studied via direct dendritic patch-clamp or via quantitative imaging of fluorescent calcium and voltage indicators (Stuart and Spruston, 2015). However, dendrites of rhythmically active interneurons in the preBöC are relatively thin (∼1-2 μm diameter) and although their branch structure is remarkably planar, they extend ∼37 μm in the parasagittal plane on average (Picardo et al., 2013), which is enough to make them difficult to target for dendritic patch-clamp. Dendrites that traverse the z-axis are also more difficult to quantitatively image since less of the branching structure can be captured within a single focal plane. Lastly, neurons in the preBöC that are visualized most easily for both whole-cell patch-clamp and imaging are found near the surface of the slice (<100 μm depth). Neural processes that traverse the z-axis near the surface are inherently prone to lesioning during slice-cutting procedures, which may compromise measurements. Limitations to the investigation of dendritic properties in acute slices have thus limited our ability to discover excitable properties of preBöC neurons beyond the peri-somatic area.

Indirect observations suggest integrative events occur on dendrites (Morgado-Valle et al., 2008; Pace and Del Negro, 2008; Pace et al., 2007a), and dendritic activity in rhythmically active neurons of the preBöC has been observed and measured (Del Negro et al., 2011). However, the manner in which integrative events in dendrites operate in a rhythmogenic context is still poorly understood. Similarly, the distribution of voltage-gated ion channels on dendrites outside the
peri-somatic area is unknown, which may additionally influence synaptic integration in the context of respiratory rhythm generation.

Dendrites likely play an important role in inspiratory burst generation due to the involvement of dendritically-localized integrating conductances (Del Negro et al., 2011; Pace and Del Negro, 2008; Pace et al., 2007a). Respiratory rhythm generation relies on synaptic excitatory input and is an emergent property of the preBötC network (Del Negro et al., 2002; Pace et al., 2007b). The group-pacemaker mechanism attempts to explain emergent network rhythm as an interplay between intrinsic membrane conductances and recurrent excitation among interconnected neurons in the preBötC. In a feed-forward process, these neurons generate a building amount of temporally-summated synaptic input. Rhythmically active neurons then amplify summated excitatory input into bursts of action potentials at the soma. The intrinsic membrane conductance responsible for the amplification of excitatory input has been identified as a calcium-activated non-specific cation current (\(I_{\text{CAN}}\)) (Beltran-Parrazal et al., 2012; Pace and Del Negro, 2008; Pace et al., 2007a), and is modulated by metabotropic glutamate receptors (mGluRs) (Pace and Del Negro, 2008). As such, recruitment of \(I_{\text{CAN}}\) likely occurs at the point of synaptic input—the dendrites. Synaptic integration occurring in dendrites can greatly influence the behavior of neurons. Voltage-dependent ion channel have been found in dendrites in nearly every mammalian neuron tested (Stuart and Spruston, 2015; Stuart et al., 2016). Depending on their exact subcellular distribution, such currents can influence the amplitude and summation of EPSPs (Magee, 2000), and in some cases support back-propagating action
potentials or calcium spiking (Kampa and Stuart, 2006; Larkum et al., 1999; Otsu et al., 2014). These phenomena allow neurons to alter their output behavior depending on the strength and timing of excitatory input (Bittner et al., 2015; Gidon and Segev, 2012; Larkum et al., 2001; Poirazi et al., 2003).

The inability to both conduct long-lasting experiments and thoroughly investigate subcellular properties might be ameliorated by the use of organotypic slice cultures. Organotypic slice cultures preserve slices of central nervous system tissue in an incubated culture environment such that the projections between regions contained in the slice and synaptic connectivity of local microcircuits are grossly retained over time. Slice cultures can be produced in one of two ways: the Gähwiler roller-tube method (Gähwiler, 1981, 1988), or the Stoppini semi-permeable membrane method (Stoppini et al., 1991). These techniques differ in their means of oxygenating brain slices. The roller-tube method depends on a slowly rotating (~5 rpm) drum that allows cover glass-mounted brain slices to be alternatively exposed to media and air. The Stoppini method utilizes semi-permeable membranes on which slices remain stationary. Membrane cultures maintain a thin layer of media at the tissue surface that allows oxygenation via passive diffusion (Figure I.2). In both methods, slices of CNS tissue can be maintained in vitro for up to eight weeks (Humpel, 2015). Thus the duration of experiments can be expanded from a single day to many weeks, including multiple recording sessions (Dong and Buonomano, 2005; Jahnsen et al., 1999; Seidl and Rubel, 2010; Yamamoto et al., 1997). The longevity of organotypic cultures permits targeted transfections and subsequent expression of vectors in one or many cells.
(Arsenault et al., 2014; Forsberg et al., 2016; Murphy and Messer, 2001; Nguyen et al., 2012; Rathenberg et al., 2003; Wickersham et al., 2007). They have also been used as models for chronic exposure treatments (Laake et al., 1999; Newell et al., 1995; Peña, 2010; Rytter et al., 2003).

One of the early applications of slice cultures was the study of developing neuronal processes (Muller et al., 1993; Zhabotinski et al., 1979; Zimmer and Gähwiler, 1987). Neurons lesioned during slicing of brain tissue either die or regenerate their neurites in cultures (Stoppini et al., 1993). As such, there is a greater likelihood that the dendrites and axons of neurons are intact when they are targeted for recording in slice cultures. The extent to which regenerated dendrites recapitulate the properties of neurons in situ or in vivo, can thus be tested. A byproduct of the culturing process is a gradual flattening-out of the slice and a concomitant increase in the clarity of the tissue (Guy et al., 2011; Humpel, 2015). As a result, the distance between synaptic partners is reduced in the z-axis, causing dendrites to become more planar. In sum, the properties of organotypic slice cultures appear to address

**Figure I. 2** The semiporous membrane organotypic culture preparation. *Left panel:* diagram of a slice placed on top of a semiporous membrane insert. Note the insert elevates the slice in the well so it does not become submerged. A thin layer of medium is maintained over the slice to prevent dessication. *Right panel:* image of a slice culture after 14 days-in-vitro.
the issues preventing long-duration experiments and the investigation of subcellular dynamics. Further, organotypic culture models have already proven successful for studying spinal cord motor rhythms (Czarnecki et al., 2008; Darbon et al., 2003; Magloire and Streit, 2009; Tscherter et al., 2001), and may thus be equally suited for patterns emerging from brainstem respiratory networks. Such precedence and increased experimental flexibility has motivated the development of a novel model of respiratory rhythmogenesis described herein.

In Chapter 1, a novel organotypic slice culture containing the preBötzinger complex is described. The spatiotemporal dynamics of rhythmic network behavior is compared to acute slices via whole-slice calcium imaging, including responses to well-established modulators of respiratory rhythm (Gray et al., 1999). We confirm that the rhythm in the culture results from the preBötC neuronal network by measuring the time course of calcium transients occurring in fluorescently labelled astrocytes versus neurons. Further, whole-cell patch-clamp recordings confirm neurons behave rhythmically as they do in acute slice preparations. This work thus affirms that organotypic cultures retain the preBötC oscillator and can be used to probe the cellular and subcellular mechanisms of rhythmogenesis.

Chapter 2 further elaborates the organotypic slice culture model of respiration and leverages its optical qualities by investigating the electroresponsive properties of neurons generating rhythm in cultures. Two behaviorally distinct classes of rhythmic neurons known to exist in acute slices are also found to persist in cultures, which bolsters our confidence in the culture model of breathing behavior. One of these neuron types is hypothesized to be rhythmogenic, the other is thought to be
premotor-related. The subcellular distribution of a transient outward potassium current (I\textsubscript{A}) is one hallmark feature of the putatively rhythmogenic neuron type and its role dendritic responsive properties is examined in both rhythmic subtypes. The results demonstrate that rhythmically active neurons featuring I\textsubscript{A} express it throughout the extent of their somatodendritic morphology. The subcellular distribution of I\textsubscript{A} alters the electrotonic compactness of neurons and could support the notion that rhythmic neurons featuring I\textsubscript{A} are most important for coordinating the onset of inspiratory burst activity.

This dissertation aims to demonstrate how current experimental obstacles in the study of respiratory rhythmogenesis can be overcome by the use of an alternative in vitro model of the behavior—the organotypic slice culture containing the preBötzinger complex. Although farther removed from the \textit{in vivo} state than other acute preparations, this culture model appears to retain the core mechanisms necessary for stable rhythmogenesis. Improved optical qualities of organotypic cultures permit detailed imaging of dendritic processes and are leveraged to demonstrate unique integrative properties in rhythmically active respiratory neurons.
REFERENCES


CHAPTER 1: Organotypic slice cultures containing the preBötzinger complex generate respiratory-like rhythms

1.1 INTRODUCTION

For nearly twenty-five years, investigators have employed acute transverse slice preparations containing the preBötzinger complex (preBötC) to probe the cellular and synaptic bases for respiratory rhythm generation in rodents (Feldman et al., 2013; Funk and Greer, 2013; Smith et al., 1991). These ‘breathing slice’ preparations retain sufficient circuitry from the respiratory medulla to spontaneously generate inspiratory-related rhythms and motor output, measurable via the hypoglossal (XII) cranial nerve root. Acute slices provide optimal access for electrophysiological, optical, and pharmacological experiments, which has helped define fundamental cellular and synaptic neural mechanisms underlying inspiratory rhythms as well as basic neuromodulatory mechanisms that influence network activity (Feldman et al., 2013; Funk and Greer, 2013; Ramirez et al., 2012).

However, acute slices remain viable and rhythmically active for approximately one day (Funk and Greer, 2013), which limits the scope of techniques that may be applied. Pharmacological assays are limited to time windows of several hours,
which prohibit studies of chronic drug effects on respiratory network function. Furthermore, experiments that involve genetic manipulation or transfection require several days for protein expression, and thus are impossible using acute slices. It would therefore be advantageous to have an in vitro preparation that retains a functional preBötzC for days or weeks, and facilitates genetic and pharmacological manipulation while retaining behaviorally relevant network activity that can be monitored and recorded with the ease offered by acute slices.

Using organotypic culturing techniques, partial slice preparations of the preBötzC have been successfully maintained over multi-week periods (Hartelt et al., 2008). These slice cultures, cut 250 µm thick, preserve some rhythmogenic function, but contain only a fraction of the preBötzC neurons normally captured in 400-700 µm thick breathing slices (Funk and Greer, 2013). Hartelt and colleagues recorded rhythmic low-frequency action potential bursts in some cells, but did not analyze whether coordinated patterns of network activity were generated in their cultures. Thus, a culture preparation that retains the preBötzC rhythmogenic core bilaterally, throughout its anterior-posterior extent, which typifies acute rhythmically active slice preparations, has not yet been described.

Here we present a novel slice preparation that retains the preBötzC bilaterally using both the Stoppini membrane method and Gähwiler roller-drum method of organotypic slice culturing (Gahwiler, 1981; Stoppini et al., 1991). This new culture preparation generates synchronized network activity bilaterally in the ventrolateral, preBötzC-related regions of the slice, analogous to rhythms in acute slice preparations. The preparation shows spontaneous rhythm over several weeks of
culturing, and flattens out, which improves the quality of optical imaging at the cellular and network level. Furthermore, the neurons are accessible for whole-cell patch clamping, meaning that both network activity and individual neurons can be selectively recorded from and perturbed using methods already established in the acute slice preparation.

1.2 RESULTS

Oscillatory activity in acute slices, brainstem slice cultures, and brainstem-cerebellar co-cultures

Bilateral oscillatory calcium activity was detected in Fluo-8 AM-loaded brainstem slices that expose the preBötC at their rostral surface (Ruangkittisakul et al., 2011, 2014), and subsequently imaged both acutely, and after 7-28 days DIV using the Stoppini culturing method (Fig. 1.1A,B). In both acute slices and cultures synchronized calcium activity was observed in an inverted V-shaped pattern with signals concentrated in dorsal (Fig. 1.1A,B, blue circles) and ventral (Fig. 1.1A,B, red circle) oscillatory groups. Acute slices showed respiratory-related rhythmic activity at 10.9 ± 4.2 bursts/min (n = 10) and slices maintained in organotypic culture for 7-22 DIV oscillated at 13.7 ± 10.6 (n = 15) bursts/min. When we co-cultured the brainstem slice with two cerebellar slice explants (placed either along the dorsolateral or ventrolateral border of the slice) then the rhythm was faster (22.4 ± 8.3 bursts/min, n = 26, Fig. 1.1C,D). Brainstem-cerebellar co-cultures (7-43 DIV) were faster than both acute slices and brainstem slice cultures, and there was no significant linear correlation, or other obvious relationship between DIV and burst frequency (not shown, linear correlation coefficient r = 0.1, n = 19).
Figure 1. Oscillatory calcium activity in three different transverse brainstem slice preparations. **A**, top panel: Cycle-triggered average of fluorescent calcium activity overlaid on brightfield image of an acute slice preparation (AS, for anatomical reference). ROIs drawn over rhythmically active areas at ventrolateral (red) and dorsomedial regions (blue). **Bottom panel**: ΔF/ΔF₀ traces of rhythmic activity. Upper/blue trace corresponds to dorsomedial ROI; lower/red trace corresponds to ventrolateral ROI. **B,C**: Same as **A**, showing calcium activity from a brainstem slice culture (BS), and a brainstem-cerebellar co-culture preparation (CC), respectively. The color calibration scale in **A** shows the colors associated with the ΔF range from minimal to maximal, and applies to all plots in figures showing ΔF images. **D**: Graphs showing burst frequency, burst duration (half-width; taken at 50% from baseline), and burst rise-time (10-90% amplitude from baseline). Error bars: Means ± SEM. *P < 0.05, **P < 0.001, ANOVA with post hoc comparison using Tukey tests. Note that the burst frequency is higher in co-culture preparations versus acute slice and brainstem slice cultures. Also, the difference in burst duration and rise time between acute slice and co-culture preparations is not significant.
Preparation type had a significant effect on frequency (F[2,48] = 8.9, p = 0.0005, Fig. 1.1D, left panel). Post hoc comparison using Bonferroni or Tukey tests indicate that co-culture oscillation frequency is significantly different than brainstem slice cultures (mean difference = 8.7 bursts/min, p = 0.008) and acute slices (mean difference = 11.5 bursts/min, p = 0.002).

Calcium transients in brainstem slice cultures were longer lasting and had a slower onset compared to acute slices or brainstem-cerebellar co-cultures. The mean burst duration of calcium activity in the ventral oscillatory group of brainstem slice cultures measured 785 ± 257 ms (n=13) compared to 367 ± 59 ms in acute slice preparations (n = 10) and 514 ± 169 ms in co-cultures (n = 26). Preparation type had a significant effect on burst duration (F[2,46] = 16.2, p = 4.7x10^{-6}, Fig. 1.1D, center panel). Post hoc comparison tests indicate that brainstem slice culture burst duration is significantly different than both acute slices (mean difference = 418 ms, p=6.1x10^{-6}) and co-cultures (mean difference = 272 ms, p=0.0002).

The rate of signal onset, measured as the 10-90% rise-time above baseline fluorescence, was slower in brainstem slice cultures (375 ± 213 ms, n = 13) compared to acute slice preparations (226 ± 43 ms, n = 10) and co-cultures (223 ± 57 ms, n = 26). Preparation type had a significant effect on signal onset (F[2,46]=7.8, p=0.001, Fig. 1.1D, right panel) and post hoc comparison tests indicate that burst rise-time brainstem slice culture is significantly different than both acute slices (mean difference = 149 ms, p=0.01) and co-cultures (mean difference = 152 ms, p = 0.001).
The signal-to-noise ratio was higher in co-cultured preparations compared to acute slices. Under similar dye-loading and imaging conditions, peak calcium signals from the center of the ventral oscillatory group measured 340% higher in magnitude in co-cultured preparations (5.5 ± 3.3 %, n = 8), and 550% higher in brainstem slice culture preparations (8.8 ± 3.1 %, n = 8) compared to acute preparations (1.6 ± 0.7 %, n = 8, (F[2,21] = 14.8, p = 9.6E-5). Active areas whose calcium activity was not normally visible in acute slices also became apparent in cultured preparations. In 53% (n = 34) of all cultured preparations, midline activity between ventral oscillatory groups was detectable.

We also prepared brainstem slices co-cultured with cerebellar explants using Roller-drum methods (Gahwiler, 1981), and these cultures (17-19 DIV) also showed spontaneous oscillatory activity, with a burst frequency of 28.6 ± 5.9 bursts/min, burst duration of 625 ± 97 ms, and 10-90% rise-time of 269 ± 29 ms. These characteristic measurements were not significantly different from similar Stoppini-type co-cultures (p = 0.09-0.16, n=5).

**Spread of activity in brainstem slice cultures and brainstem-cerebellar co-cultures**

Both brainstem slice cultures and brainstem-cerebellar co-cultures display oscillatory calcium burst patterns that propagate ipsilaterally, travelling from their point of initiation in the ventrolateral area toward the dorsomedial border at varying velocities (Fig. 1.2A). When imaged at 129 Hz, the signal in brainstem-cerebellar co-cultures propagated 275 % as fast (0.022 ± 0.007 m/s, n = 8) as brainstem slice cultures (0.008 ± 0.005 m/s, n = 9, p<0.001; Fig. 1.2B). In each slice there was a
detectable latency in the signal rise between contralateral ventral oscillatory groups. In every case, for both brainstem slice cultures and brainstem-cerebellar co-cultures, calcium signals took longer to propagate from ventral to dorsal regions than to contralaterally equivalent regions, where the activity appeared within 1-2 frames, which made it too uncertain to calculate a precise velocity or possible dominant or alternating initiation site (n = 7). Thus, regardless of preparation type, synchronized bursts first occurred bilaterally between contralateral ventral oscillatory groups and then propagated more slowly in the dorsomedial direction.

**Ventral oscillatory group activity in multiple neurons and calcium transients in dendritic profiles**

In acute slice preparations, calcium and voltage dye fluorescence imaging often appears as diffuse regional fluorescence due to light scattering in the tissue, or instead captures only a subset of rhythmically active neurons within the field of

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**Figure 1.2** Propagation of calcium activity in brainstem slice cultures and brainstem-cerebellar co-cultures. **A:** Time series Z-projection of maximum fluorescent calcium activity in a brainstem-cerebellar co-culture. The heat map overlay shows threshold signal increases from low (red) to high latencies (blue) during a single rhythmic burst acquired at 129 frames/s. **B:** Average velocity of signal propagation in brainstem slice cultures (BC) versus brainstem-cerebellar co-cultures (CC). Error bars: Means ± SEM. **P < 0.001, Student’s t-test."
scattering at locations deep within the slice (Del Negro et al., 2011; Funk and Greer, 2013; Katona et al., 2011). One advantage of culturing brain slices is the reduced thickness compared to acute slices (Stoppini et al., 1991). Using the Stoppini culturing method, we reduced the thickness of the brainstem slices cultured here to 104 ± 31 μm (measured in sagittal sections of Epon-embedded 7-31 DIV oscillating cultures, n = 5) from a thickness of 400 μm in the acute slices at
0 DIV. The dorsal-to-ventral distance at the slice midline increased to 1943 ± 183 µm compared to 1490 ± 74 µm in acute slices acquired from age-matched controls (P2.5, p < 0.05, n = 5). Thus, the cultures flatten out and the reduced thickness made it possible to record synchronized calcium transients from hundreds of neurons in the same focal plane (Fig. 1.3A,B). At higher magnification both somatic and proximal (~100 µm) dendritic activity could be recorded in Fluo-8 AM-loaded neurons (Fig. 1.3C).

**Retrograde biocytin labeling in Roller-drum cultures**

The preBötC contains commissural interneurons, which synchronize bilateral halves of the preBötC, as well as interneurons that project ipsilaterally to premotor neurons in the intermediate reticular formation, as well as hypoglossal motor neuron pools (Wang et al., 2014). If the inverted V-shaped pattern of rhythmic calcium activity in cultures corresponds to that seen in acute slices, then similar axon projections ought to be maintained in the cultured preparation. To test this prediction, we electroporated 1% biocytin using patch pipettes placed in the ventral oscillatory group of Roller-drum cultures (n = 3). Retrogradely labeled neurons were located around the injection site, somatas and projecting fibres were found in the ipsilateral dorsal part the culture, the midline, and in the contralateral ventrolateral part of the cultures (Fig. 1.4B,C). The somatic positions of the labeled neurons and fibres corresponded to oscillatory regions recorded before electroporation (Fig. 1.4A,B). The biocytin labeling suggests that the pattern of activity in cultures is attributable to an underlying bilateral network of neurons in
Biocytin-labeled fibers entering the cerebellar explant, terminating in bouton-like structures were detected in one slice co-culture (Fig. 1.4B,D), indicating a degree of synaptic interconnectivity between the brainstem and cerebellar neurons.

**Calcium activity in GFAP+ labeled cells**

Oscillatory calcium transients have been reported in astrocytes in sync with neighboring preBötC neurons (Oku et al., 2015). We tested for rhythmic glial activity in brainstem-cerebellar co-cultures prepared from transgenic mice that express EGFP coupled to glial fibrillary acidic protein (Fig. 1.5). However, oscillatory calcium fluorescence was undetectable in 98% (n = 288, 10 cultures) of
the recorded GFAP+ cells (Fig. 1.5A). A few EGFP-expressing astrocytes (n = 5) generated one-time calcium transients independently of calcium transients in neighboring oscillating neurons (Fig. 1.5B, n = 4). The mean rise-time of these transient events (810 ± 235 ms, n = 5) exceeded the length of calcium transients...
in neighboring rhythmic neurons (393 ± 66 ms; n = 119 neurons, across 5 cultures, p=0.0051). Half-width duration of transients in GFAP-cells (1028 ± 124 ms) was 24% longer than rhythmic neurons (866 ± 31 ms; n = 119 cells, across 5 cultures, p = 0.022).

**Whole-cell electrophysiology**

To assess whether neuronal behavior in cultures resembles preBötC neurons in acute slice preparations (Funk and Greer, 2013), we recorded the voltage trajectory in rhythmic neurons from culture preparations. First, using calcium imaging, rhythmically active cells within the ventral oscillatory group were targeted for whole-cell patch-clamp recordings (Fig. 1.6A, n = 10). Rhythmic drive potentials (343 ± 207 ms duration, n = 7) that reflect underlying network activity were recorded at typical membrane potential (overriding action potentials were removed via bandpass filtering), and at hyperpolarized potentials using negative bias current to uncover non-linear membrane behavior indicative of recruitment of active conductances (Fig. 1.6B). The amplitude of the underlying drive potential was 185% larger in rhythmic neurons held at close to resting membrane potential (19.0 ± 6.6 mV; n = 7), than in neurons held at membrane potentials below the threshold of action potential generation during the burst (10.3 ± 3.7 mV; n = 5, p < 0.05).

Finally, we tested pharmacological modulation of burst frequency by neuropeptides. From a control frequency of 24.4 ± 6.4 bursts/min, the NK1-receptor agonist Substance P (SP, 500 nM) sped up the frequency by 134% to 32.7 ± 9.3 bursts/min (p<0.05), and adding the µ-opioid agonist DAMGO (1 µM)
on top of SP slowed the frequency by 50% to 16.5 ± 8.7 bursts/min (p<0.01, n= 6 cumulative dosings, Fig. 1.6C).

Figure 1.6 Whole-cell patch clamp recordings from a rhythmically active neuron in the ventral oscillatory group of a brainstem-cerebellar co-culture after 7 DIV. A, Top trace: normal bursting activity at resting Vm with zero current bias applied. Bottom trace, Rhythmic drive potential at a hyperpolarized potential, after negative bias (-0.1 nA) applied. B: Overlaid cycle-triggered, and action potential-filtered average traces of the burst events occurring in A and B. Note that the underlying burst envelope is larger in amplitude at resting Vm levels. C. Modulation of burst frequency by 500 nM Substance P (SP), and 500 nM SP + 1 µM DAMGO. Error bars: Means ± SEM. *p < 0.05, **p < 0.001, n = 6, paired sample t-Test, control versus SP, and SP versus SP plus DAMGO.
1.3 DISCUSSION

Acute slice preparations containing the preBötC survive in organotypic culture conditions and maintain patterned rhythmic calcium activity, which we attribute to underlying neuronal inspiratory-like drive potentials and spike bursts in constituent preBötC interneurons. Bursts in culture begin with fast bilateral co-activation in ventrolateral regions of the slice and then propagate ipsilaterally to the dorsomedial regions of the culture preparations. The inverted V-shaped activity pattern was consistent; it did not depend on the presence of cerebellar explants. The ipsilateral ventral-to-dorsal propagation velocity was on the order of 0.02 m/s in brainstem-cerebellar slice cultures. This velocity is roughly 10 times slower than a published value of 0.24 m/s in the commissural fiber tracts connecting bilateral preBötC in rhythmically active acute slices of neonatal rats using using voltage-imaging (Koshiya et al., 2014). Thus, the slice cultures appear to generate patterned activity that spreads dorsally slower than the normal velocity of fine axons abundant in the neonatal central nervous system (Sternberger et al., 1979), which may imply that the activity is shaped by local synaptic processes in networks of neurons. The pattern also overlaps the expression of cells—in the transverse plane of the brainstem and cervical spinal cord—derived from the homeodomain transcription factor, Dbx1, which gives rise to rhythmogenic interneurons that comprise the preBötC (Bouvier et al., 2010; Gray et al., 2010; Picardo et al., 2013) and also play a role in premotor drive transmission (Wang et al. 2014).

In some recordings we also observed midline activity, which may reflect raphe neurons that interconnect with the preBötC and increase firing frequency in
response to rhythmic preBötC input (Ptak et al., 2009). Retrograde biocytin labeling from the ventral oscillatory group reveals projections in both the contralateral ventral oscillatory group and dorsomedial regions, similar to that obtained via in vivo fluoro gold injection into the preBötC (Koshiya et al., 2014). Retrogradely labeled neurons were also found in the midline, further suggesting the involvement of raphe neurons. Thus it appears unlikely that rhythmic calcium changes in the slice occur due to spontaneous calcium oscillations, but rather they resemble the neuronal behavior found in acute slices, similar in both dynamics and patterning, and occurring over anatomical regions that correspond to known areas of activity, and to locations of cells vital for production of the respiratory rhythm. The presence of cerebellar explants co-cultured with preBötC slices bolsters the rhythm, causing it to oscillate faster than acute preBötC preparations, but with similar burst duration and 10-90% rise-times. Without the addition of cerebellar explants, brainstem cultures oscillate at the same frequency as acute preparations but individual bursts have a slower onset and offset—increased burst duration and rise time. Although the specific roles of cerebellar explants that promote preBötC-like rhythmic activity in cultures are unknown, spontaneous oscillatory activity in cerebellar granule cells may contribute to the overall excitability of the slice (Apuschkin et al., 2013; De Zeeuw et al., 2008), especially if co-culturing allows granule cell axon entry into the preBötC and facilitates excitatory synaptic connections with preBötC interneurons of the ventral oscillatory group. Since preBötC-like rhythmic behavior occurred in the absence of cerebellar explants, but that these rhythms were faster with cerebellar co-cultures, we conclude that
cerebellar circuits augment excitability in the cultured preBötC but do not contribute to the cellular mechanisms underlying rhythm generation.

Due to gradual thinning of the slice and consequent reduction of light scattering in the tissue over time, the brainstem slice culture preparation allows for higher spatial resolution when imaging at both the cellular and network level and greater amplitude of fluorescent signals. This could be advantageous for analyzing larger portions of rhythmically active cell populations in the preBötC and adjacent premotor areas than has previously been possible in acute slices. For example, the culture preparation we characterize here may facilitate real-time visualization and perturbation of individual burst percolation across the rhythmogenic core and into premotor areas. Our preparation makes it feasible to stimulate and record large populations (~100s of neurons), since neighboring somata that would otherwise be spread through the Z-axis become coplanar (or nearly so for practical purposes). These conditions also simplify imaging requirements: population-level recordings could be acquired using wide-field excitation. Additionally, subcellular sections of neurons (e.g. dendritic processes) can be visualized at higher magnifications. Therefore, it is more likely that several hundred micron regions of dendritic processes can be captured in a single focal plane, and thus intercellular signal propagation arising from recurrent excitation along the dendritic arborization of rhythmic neurons, and its impact on burst generation, might also be more readily visualized.

Astrocytes have been implicated in respiratory rhythm generation (Hartel et al., 2009; Hulsmann et al., 2000; Oku et al., 2015; Schnell et al., 2011). We tested for
the presence of calcium oscillations in the astrocytes of brainstem-cerebellar co-cultures. We were able to image a large number of cells simultaneously using conventional wide-field illumination microscopy. We used a bath-applied red-shifted calcium indicator, despite an inherently reduced fluorescence intensity compared to brighter green dyes (e.g. OGB1, Fluo-8), because of the flattened state and reduced tortuosity in the culture preparation. There was no apparent pre-inspiratory activity or oscillatory activity in 288 recorded GFAP\(^+\) cells. However, we cannot entirely rule out low amplitude calcium oscillations. Previous recordings from astrocytes in preBötC brainstem slices employed the high-affinity indicator Oregon Green BAPTA 1 (K\(_d\) = 170nM) whereas we used the medium-affinity indicator Asante Calcium Red (K\(_d\) = 400nM). Previously reported astrocytic fluctuations that were of small amplitude (<1\% \(\Delta F/F_0\)) and only visible after filtering (Oku et al., 2015). It is therefore plausible that we failed to detect astrocytic fluctuations due to experimental limitations related to our green GFAP reporter strain coupled with a red calcium dye of limited sensitivity. Further, the tight proximity of cell layers in culture conveniently allows many cells to be recorded simultaneously with less light scattering due to tissue thickness, but the signal is likewise prone to contamination by neuronal activity that is marginally out of focus yet still of a greater magnitude than any glial signals, which may be especially true at higher magnification. Finally, it is possible that astrocytes in this case were inadequately labeled due to loading feasibility of the indicator dye, or that they simply behave differently in a culture environment.
Whole-cell electrophysiology data show neurons in the ventral oscillatory group of brainstem-cerebellar co-cultures whose behavior appears similar to previously described rhythmic neurons in the preBötC of acute slices (Funk and Greer, 2013; Ramirez et al., 2012). These neurons receive periodic inspiratory drive potentials that generally produce bursts of action potentials with an underlying burst envelope that is larger at rest than at hyperpolarized potentials (Fig. 1.6B), suggesting that voltage- or calcium-dependent conductances amplify the synaptic drive (Ramirez et al., 2012). The drive potentials had a duration of ~350 ms, which compares to the ~870 ms half-amplitude duration of the calcium signal in individual neurons, illustrating that the shape and time course of the calcium signal also depend on cellular calcium kinetics. Taken in conjunction with calcium imaging data and retrograde tracing with biocytin, we conclude that organotypic cultures retain a rhythmically active ventral network that corresponds to the preBötC. Its activity pattern spans dorsally into regions such as the intermediate reticular formation, the nucleus tractor solitarius, and the XII nucleus, which are associated with premotor and motor circuits that serve respiration. Midline activity also appears to correspond with the raphe obscurus (Ptak et al., 2009). Decreased tissue thickness and consequently the ability to capture many active neurons in the same focal plane with significantly reduced fluorescence scattering likely accounts for the increase in signal amplitude in cultures (both brainstem slice cultures and brainstem-cerebellar co-cultures) compared to acute slices. However, the existence of rhythmic neurons in the ventrolateral slice culture, and adjacent dorsomedial regions, does not necessarily guarantee that these neurons
correspond one-to-one to preBötC interneurons and respiratory premotor or motor neurons, or that they have not modified their behavior or physiology in some way. A degree of synaptic remodeling certainly occurs over time in vitro—we can see promiscuous fibers projecting between cerebellar and brainstem explants via retrograde biocytin tracing. Thus, we cannot exclude that circuits not found in the intact animal or unnatural strengthening of existing circuits develop over time in the cultures, which certainly should be taken into consideration when using this preparation. However, general synaptic connectivity between contralateral and dorsomedial motor regions of the slice appears to be preserved and, more importantly, the calcium activity in preBötC cultures reflects a consistent rhythmic pattern—initiation in the ventral oscillatory group followed by fast bilateral synchronization and subsequent propagation of signal to dorsomedial regions. Pharmacological modulation of the rhythm using agonists for NK1 and µ-opioid receptors also mimics responses seen in acute slice (Gray et al., 1999), demonstrating that essential peptidergic modulating systems are intact in the cultures, at least at the postsynaptic level.

Thus, the preBötC slice culture preparation could be a useful model of respiratory rhythm generation that approximates the already widely used acute slice preparation and is amenable for optical and electrical stimulation and recording. The longevity of the culture preparation will facilitate molecular biological experiments and techniques, as well as any protocol that requires pharmacological perturbations lasting multiple days or weeks.
1.4 METHODS

Ethical approval
The Department of Experimental Medicine at the Panum Institute approved all experiments and procedures according to protocols laid out by Danish Ministry of Justice and the Danish National Committee for Ethics in Animal Research.

Organotypic slice cultures
US Naval Medical Research Institute (NMRI) mice post-natal ages 0.5 to 5.5 days were anesthetized with isoflurane and immediately dissected in sterile-filtered chilled artificial cerebrospinal fluid (ACSF) containing (in mM): 184 glycerol, 2.5 KCl, 1.2 NaH2PO4, 30 NaHCO3, 20 HEPES, 25 D-glucose, 5 sodium ascorbate, 2 thiourea, 3 sodium pyruvate, 10 MgSO4, 0.5 CaCl2; pH 7.3, equilibrated by bubbling with carbogen (95% O2 and 5% CO2). We cut transverse slices of the brainstem, 400 μm in thickness, at the level of the preBötC on a vibratome (Thermo Scientific Microm 650V, Waltham MA, USA) as described by newborn mouse atlases of the brainstem (Ruangkittisakul et al., 2011, 2014). Additionally, we often retained 400-μm-thick parasagittal slices of the cerebellum from the same mouse brain for co-culturing. For Stoppini-type cultures, transverse brainstem slices were placed onto semi-porous culture well inserts (Millipore PIC03050, Darmstadt, Germany) with cerebellar explants co-cultured along their dorsolateral borders. Mounted preparations were maintained in sterile-filtered organotypic culture media containing: 50% Eagle’s MEM with Earle’s Salts, 25% Hank’s balanced salt solution, 25% heat-inactivated horse serum, 2 mM glutamine, 200 U/ml penicillin, 5 μg/ml streptomycin, 10 mM HEPES and an additional 3.6 mM D-glucose. The
cultures were treated with 10 µM MK-801, an NMDA receptor antagonist, for the first three days in vitro. Fresh culture media was supplied every 48 hours thereafter until experimentation. These slices were then kept in a sterile humidified incubator at 35° C and atmospheric CO₂ concentrations during incubation.

We prepared some cultures according to the roller-drum technique, wherein the slices were placed on a coverslip embedded in citrated chicken plasma (Sigma-Aldrich, St. Louis, MO, USA) that was then coagulated by adding a drop of 100-150 U/µL thrombin (Sigma-Aldrich). The cultures were kept in sealed plastic tubes (flat bottom, NUNC Thermo Scientific, Waltham MA, USA) and placed in a rotating roller-drum (15 revolutions/h) at 35° C in humidified air. All procedures were performed in a sterile work environment within a Class 1 biosafety cabinet. The success rate in obtaining oscillating cultures depended in particular on the dissection experience and skill of experimenter, and approached 100% after ~3 months of training.

The thickness of the cultures was measured in counterstained sections using a Zeiss Axiolab microscope: The culture was drop fixed overnight in 4% paraformaldehyde, washed in Sørensen’s Phosphate Buffer (0.1 M), dehydrated with increasing ethanol, embedded in Epon (EMBed-812, Electron Microscopy Sciences, PA, USA), cut in 3 µm sections at a 90 degrees angle relative to the surface on a microtome, counter stained with 0.1% Toluidine (Sigma-Aldrich), and mounted with the mounting medium, Eukitt (Sigma-Aldrich).
GFAP+ mice

Mice expressing enhanced green fluorescent protein (EGFP) under the promoter of glial fibrillary acidic protein [GFAP; line Tgn(hgFAPEGEP) GFEC 335] were kindly provided by Professor Frank Kirchhoff at the University of Saarland in Homburg, Germany (Lalo et al., 2006).

Electroporation

Glass micropipettes were pulled from filamenteed capillary glass (outer diameter 1.5 mm, inner diameter 0.86 mm, Harvard Apparatus, Holliston, MA, USA) using a PUL-100 micropipette puller (World Precision Instruments, Sarasota, FL, USA), giving a tip diameter of 1-3 µm. Pipettes were filled with 1% biocytin, 1 M KCl, 0.126 M Tris, placed in the slice culture, and positive current was applied in 50 ms pulses of 200 µA at 2 Hz for 10 min. Current pulses were generated by a stimulus isolation unit gated by a waveform generator (IsoFlex and Master-8, A.M.P.I., Jerusalem, Israel).

Calcium imaging, and peptidergic compounds

Loading solutions for membrane-permeable calcium indicators were prepared by combining 30 µl of a 10 mM stock solution containing either Fluo-8 AM in DMSO (AAT Bioquest, Sunnyvale, CA) or Asante Calcium Red AM in DMSO (TEFLabs, Austin, TX, USA) with 3.5 µl cremophore EL (Fluka, St. Louis, MO, USA) and 7.5 µL of 20% pluronic acid in DMSO (AAT Bioquest, Sunnyvale, CA, USA). This 41 µl of dye solution was then dissolved in 1.5 ml of a standard artificial cerebrospinal fluid solution (ACSF) containing 10 µM MK-571 (an ATP-binding cassette transporter blocker that improves calcium indicator dye uptake (Manzini et al.,
2008). The ACSF consisted of (in mM): 129 NaCl, 3 KCl, 25 NaHCO$_3$, 5 KH$_2$PO$_4$, 30 D-glucose, 0.7 CaCl$_2$, 0.4 MgSO$_4$, 100 mM D-mannitol, which was aerated by bubbling with carbogen (95% O$_2$ and 5% CO$_2$) at room temperature. The final Fluo-8 AM (or Asante Calcium Red) concentration was 20 µM. Slice cultures were submerged in bubbled loading solution for 30-60 minutes before recording.

Fluorescent calcium activity was recorded in the slice culture using a stereo microscope (Leica MZ16 FA, Wetzlar, Germany) 30 min after the preparation was placed in a 2-ml recording chamber at 29°C, and perfused at 2 ml/min with preheated oxygenated ACSF. Light-emitting diodes (LEDs, HLV series, CCS LED Spotlight, Kyoto, Japan) and a metal-halide light source (Leica EL6000) illuminated the preparation. Appropriate filter sets for green fluorescence (Leica GFP3: excitation band-pass 450-490 nm and emission band-pass 500-575 nm) were inserted in the light path for fluorescence imaging. Time series acquisition was performed with an electron-multiplying charge-coupled device (EMCCD) camera (LucaEM S DL-658M, Andor Technology, Belfast, United Kingdom) at 10-129 Hz, controlled by SOLIS software (Andor Technology).

For cellular-level fluorescence detection, a metal halide light source PhotoFluor II (89North, Burlington, USA) or a LED light source, M470L2 (Thorlabs, Newton, NJ, USA) was coupled to a fixed-stage upright microscope via a liquid light guide and appropriate optical filters (for Fluo-8 AM, a modified Olympus U-MWIB set: excitation 457-487 nm, dichroic mirror 505 nm, emission 515-550 nm. For Asante Calcium Red AM, an Olympus U-MWIG2 set: excitation 520-550 nm, dichroic mirror 565 nm, emission BA580IF nm). Images were captured by an sCMOS
camera (Neo DC-152Q, Andor Technology) controlled by the SOLIS software (Andor Technology). Imaging protocols employed a 10x (NA 0.3), 20x (NA 0.5), 40x (NA 0.8), and 63x (NA 0.95) water immersion objectives. Time series images were acquired at 10–50 frames/s.

The NK1-receptor agonist Substance P acetate salt hydrate (SP), and µ-opioid agonist \([D-Ala^2, N-Me-Phe^4, Gly^5-ol]-Enkephalin\) acetate salt (DAMGO, Sigma-Aldrich) were dissolved in water and further diluted in ASCF to give final concentrations of 500 nM and 1 µM respectively. The effect of Substance P and DAMGO on burst frequency was tested in cumulative dosing measuring burst frequency over 20 s image stack in the control, 10 min of bath applied Substance P, and 10 min of Substance P plus DAMGO.

**Whole-cell patch clamp**

Patch recordings employed an AxoClamp 2B amplifier (Molecular Devices, Sunnyvale, CA, USA) and ROE-200 micromanipulators (Sutter Instruments, Novato, CA, USA). Glass patch pipettes were pulled as described above to a tip resistance of 9-11 MΩ. Pipettes were loaded with a patch solution containing (in mM): 165 potassium D-gluconate, 10 NaCl, 0.5 MgCl₂, 10 HEPES, 0.4 GTP, 4 ATP, 0.5 EGTA, pH 7.3, and then visually guided to target cells using a fixed-stage upright microscope (modified Olympus BX51) with a 63x (NA 0.95) objective. Data were digitally acquired at 20 kHz. Rhythmically active neurons were first recorded with the minimum necessary current bias to achieve stable activity with little spontaneous firing during inter-burst intervals.
Analysis and statistics

Optical and electrophysiological data were analyzed offline using Igor Pro v. 6.36 (Wavemetrics, Lake Oswego, USA), Clampex 10.3 (Molecular Devices), and ImageJ v.1.49 (Schneider et al., 2012). Change in fluorescence over baseline fluorescence intensity ($\Delta F/F_0$) was calculated using a moving Z-projection that finds minimum values of fluorescence across consecutive 1-s time windows throughout an image stack. The 1-s running average, uniquely calculated for each individual frame, was subtracted from it, throughout the time series to isolate fast calcium fluctuations from background fluorescence. Image stacks were then Kalman filtered to reduce noise and pseudo-colored (‘rainbow’ RGB look-up-table, red: maximum values of $\Delta F$, green: medium values of $\Delta F$, blue: minimum values of $\Delta F$). Pixels were binned (mean of adjacent 2x2-10x10 points) and brightness and contrast were enhanced using ImageJ. In some experiments regions of interest (ROIs) were defined and average $\Delta F$ values within a given ROI were plotted versus time. Unless otherwise stated, statistical values are given as mean ± S.D. Student’s t-test was used for statistical comparisons of two sample populations and ANOVA when more than two sample populations are compared. All statistical tests were performed using Origin 2015 (OriginLab Corp., Northampton, MA, USA).
1.5 REFERENCES


CHAPTER 2: Dendrites of rhythmically active neurons in the preBötzinger complex contain an \( I_A \)-like potassium current

2.1 INTRODUCTION

Breathing is essential for homeostasis. Therefore, the physiological properties of neural circuits that generate and control respiratory rhythm are of longstanding interest. The preBötzinger complex (preBötC) of the ventral medulla contains a network of excitatory interneurons that generate the rhythm for inspiratory breathing movements (Feldman and Del Negro, 2006; Feldman et al., 2013; Smith et al., 1991). preBötC neurons can be dichotomously subdivided into two classes, which differ with respect to membrane potential trajectory during the respiratory cycle and electroresponsive properties (Picardo et al., 2013; Rekling et al., 1996). During rhythmic activity, type-1 preBötC neurons integrate synaptic drive and exhibit preinspiratory depolarization ~400 ms prior to inspiratory bursts. Type-2 neurons integrate synaptic as well, but it occurs later in the cycle, and type-2 neurons exhibit preinspiratory depolarization ~200 ms prior to inspiratory bursts (Rekling et al., 1996). Early preinspiratory activity in type-1 neurons suggests that they may initiate recurrent excitation that leads inexorably to the
synchronized onset of inspiratory burst and thus be rhythmogenic (Rekling et al., 1996; Smith et al., 1990).

The biophysical basis for the synchronized and coordinated onset of activity across the rhythmic neuronal population (i.e. orderly network recruitment) initiated by type-1 neurons has yet to be fully explained, but may be related to the presence of a transient outward $I_A$-like current, a defining feature of type-1 neurons (Rekling et al., 1996). Transient outward potassium currents (i.e., $A$-currents, $I_A$) exhibit a range of voltage-dependent and kinetic properties (Birnbaum et al., 2004). In the preBötC, $I_A$ activates at subthreshold membrane potentials and fully inactivates at membrane potentials above -40 mV (Hayes et al., 2008). Pharmacological inhibition of $I_A$ in the preBötC results in spurious burst generation at both the cellular and network levels, which the authors attributed to disorderly recruitment of rhythmogenic preBötC neurons during recurrent excitation that precedes the inspiratory burst in each cycle of the rhythm (Hayes et al., 2008).

Measured via dendritic patch-clamp and Ca$^{2+}$ imaging, $I_A$ is expressed at a high density on the dendrites of hippocampal pyramidal cells and cerebellar Purkinje cells, where it acts locally to inhibit excitatory responses like Ca$^{2+}$ spikes (Hoffman et al., 1997; Kampa and Stuart, 2006; Otsu et al., 2014). Dendritic $I_A$ limits the effect of excitatory inputs that are sparse or transient (or both). Conversely, dendritic $I_A$ promotes robust post-synaptic responses to temporally summated excitatory input, which is long-lasting enough to inactivate $I_A$ and thus
exert a more profound influence on membrane potential trajectory (Magee et al., 1998).

The amplitude of the drive potential underlying inspiratory bursts in rhythmic preBötz neurons depends on a calcium-activated non-specific cation current ($I_{\text{CAN}}$), which can be activated by $\text{Ca}^{2+}$ influx via voltage-gated channels or via intracellular $\text{Ca}^{2+}$ release linked to group I metabotropic glutamate receptors (mGluRs) (Pace et al., 2007). Simultaneous dendritic application of AMPA and group I mGluR agonists, but not AMPA alone, amplifies excitatory input recorded at the soma (Pace and Del Negro, 2008), suggesting that excitatory input is amplified by intrinsic conductances on dendrites.

Temporally sparse spontaneous synaptic activity during the interburst interval could cause spurious burst generation, if such activity could also evoke $I_{\text{CAN}}$. However, a mechanism that selectively inhibits sparse synaptic inputs could prevent dendritic amplification, thus avoiding spurious bursts during the interburst interval. Fast-activating voltage-dependent outward current on dendrites can shunt sparse excitatory input, and might explain the aforementioned disordered behavior seen during network-wide blockade of $I_A$ in rhythmically active slices. Thus, we hypothesize that $I_A$ is expressed on dendrites in type-1 neurons, in addition to being somatically expressed (Hayes et al., 2008). Here we utilize an organotypic slice culture containing the preBötz to investigate the contribution of $I_A$ in rhythmically active preBötz neurons to influence transient depolarization along dendrites and find that loss of dendritically localized $I_A$ significantly increases the amplitude of voltage-sensitive dendritic $\text{Ca}^{2+}$ transients. These
phenomena demonstrate a possible mechanism by which dendritic $I_A$ might filter synaptic input to type-1 neurons, blocking sparse input yet permitting temporally and spatially summating inputs during the preinspiratory phase, which can inactivate $I_A$, to exert a more profound effect.

2.2 RESULTS

Establishment of two rhythmic neuron classes in organotypic cultures of the preBötC

Neurons in organotypic slice cultures containing the preBötC retain rhythmic respiratory related behavior analogous to that seen in acute slice preparations and brainstem en bloc recordings (Forsberg et al., 2016; Funk and Greer, 2013; Phillips et al., 2016). However, it is not yet known whether rhythmically active preBötC neurons in culture retain the dichotomous membrane properties classified as type-1 and type-2 first characterized in acute slice recordings (Rekling et al., 1996). Type-1 and type-2 neurons are segregated based on membrane voltage trajectory during rhythmic behavior (e.g., pre-inspiratory activity) and the presence of either a transient outward current, $I_A$, or a hyperpolarization-activated inward current, $I_h$. Type-1 neurons prominently feature $I_A$, lack $I_h$, and display a ramp-like increase in membrane potential, known as pre-inspiratory activity, beginning ~400 ms prior to the inspiratory burst. Conversely, type-2 neurons lack $I_A$, express $I_h$, and display more latent pre-inspiratory activity ~200 ms prior to the inspiratory burst (Rekling et al., 1996). Motor nerve rootlets typically deteriorate in slice cultures after 6 to 7 days in vitro.
(DIV), so here type-1 and type-2 neurons were differentiated based on the presence of either $I_A$ or $I_h$, but not preinspiratory latency.

Figure 2.1 Electroresponsive properties of oscillating type-1 and type-2 neurons. A: Spontaneous oscillatory burst activity in two neurons recorded in current camp mode. Note that the burst in the left-most neuron shows afterhyperpolarizations following the bursts. B: Hyperpolarizing and depolarizing square current pulses from around resting $V_m$, and from a slightly hyperpolarized membrane potential give rise to two distinctive electroresponsive responses in the two neurons. The type-1 neuron show delayed excitation (arrow, top right trace), and the type-2 neuron show sag-rebound potentials (arrows, lower left). Voltage traces are cycle-triggered averages of 5-10 sweeps, which truncates action potentials but retains the form of delayed excitation and 'sag' potentials.
Here we obtained whole-cell recordings from rhythmically active neurons in organotypic cultures containing the preBötc and tested the presence of both $I_A$ and $I_h$ in current clamp (Fig. 2.1, $n = 42$). Rhythmic activity was first recorded using the minimum amount of negative holding current to inhibit spontaneous action potentials between rhythmic bursts (Fig. 2.1A, 0 to -0.15 nA). Among all rhythmically active neurons, the mean burst interval was $5.4 \pm 3.2$ s, and the mean burst duration was $405 \pm 134$ ms ($n = 42$).

The presence of $I_A$ was determined by first hyperpolarizing neurons with negative holding current to a baseline membrane potential ($V_m$) between -70 mV and -80 mV, which fully deinactivates $I_A$ (Hayes et al., 2008). Square-wave positive current pulses of 400 ms duration were then delivered to evoke repetitive firing of action potentials. Neurons with $I_A$ display a delay in membrane depolarization lasting 100-200 ms before firing repetitively, whereas neurons without $I_A$ discharge action potentials throughout the duration of the current pulse without any notable delay exceeding the membrane time constant (Fig. 2.1B shows cycle-triggered averages of many sweeps to demonstrate the repeatability of delayed excitation or the lack thereof in type-1 and type-2 preBötc neurons).

The presence of $I_h$ was determined by setting baseline $V_m$ between -40 and -50 mV and delivering 400 ms negative current pulses of sufficient amplitude to hyperpolarize the neuron to -70 to -90 mV, which is sufficient to evoke $I_h$ if it is expressed by the neuron. Neurons with $I_h$ exhibit a ‘sag’ depolarization of ~10 mV after being transiently hyperpolarized as well as a post-inhibitory rebound after the negative current pulse terminates (Fig. 2.1B shows cycle-triggered...
averages of many sweeps to demonstrate the repeatability of ‘sag’ or the lack thereof in type-1 and type-2 preBötC neurons).

Among all recorded neurons we found 57% (n = 24) displayed type-1 properties of delayed excitation (i.e., I_A) and lack of sag potential (i.e., no I_h), while 31% (n = 13) displayed type-2 properties of sag potentials (I_h) and lack of delayed excitation (no I_A). Of the remaining neurons, 7% (n = 3) showed both sag potential and delayed excitation, while 5% (n = 2) displayed neither I_A nor I_h. These results are in line with the distribution of respiratory neuron classes in acute slices (Picardo et al., 2013; Rekling et al., 1996). If we consider the null hypothesis to be that there is no relationship governing the expression of I_A and I_h in respiratory neurons, then the allotment of recorded neurons into type-1 and type-2 phenotypes is unlikely to have occurred by random chance (Fisher’s exact test, p<0.0001), suggesting that the dichotomous electrophysiological properties of rhythmically active type-1 and type-2 neurons observed in the preBötC from acute slices, also persists in culture.

**Voltage ramps during blockade of I_A increase electrotonic compactness of type-1 neurons**

To determine whether I_A might actively inhibit the spread of voltage transients along dendrites in type-1 neurons, we next performed simultaneous single-electrode voltage clamp (SEVC) and fluorescent Ca^{2+}-imaging of rhythmically active neurons dialyzed with the fluorescent Ca^{2+} indicator, Fluo-8L (Fig. 2.2). This allowed us to track the relative amplitude of voltage changes at distal
Figure 2. 2 Bath applied 4-AP increases dendritic Ca\textsuperscript{2+} transients in response to ramp depolarizations. 

**A:** Voltage clamp traces, with TTX (1 µM) in the perfusate, before and after adding 2 mM 4-AP to the bath. Left-most neuron shows a transient outward current, which is blocked by 4-AP thereby classifying the neuron as type-1. Right-most neuron shows no evidence of a transient outward current, classifying the neuron as a type-2. 

**B:** Ca\textsuperscript{2+} transients in proximal and distal dendritic compartments in the two neurons in response to a 150 ms voltage ramp (-80 to 20 mV), and associated voltage and current traces. Responses before and after adding 2 mM 4-AP to the bath are overlaid. *Blue lines:* Proximal dendritic compartments. *Green lines:* Distal dendritic compartments. 

**C:** Live morphology of the two neurons with thresholded dendritic Ca\textsuperscript{2+} transient amplitude overlaid in red. Red indicates Ca\textsuperscript{2+} transient amplitudes ($\Delta F/F_0$) above 100%. 

**D:** Group data expressing the relative Ca\textsuperscript{2+}-transient amplitude after 4-AP-from all type-1 (n=6) and type-2 (n=5) neurons.
dendritic compartments using voltage-sensitive Ca\textsuperscript{2+} influx as a surrogate for direct measurements of membrane potential.

Since voltage clamping dendrites inherently suffers from a lack of space clamp as a function of distance from the recording pipette, changes in membrane potential enforced at the soma via SEVC would be attenuated at distal dendritic locations. The exact amount of that attenuation depends on passive cable properties (e.g. length, diameter, branch order) and density of voltage-gated conductances (Bar-Yehuda and Korngreen, 2008), but space-clamp error must be factored into data analysis and interpretation.

Neurons were first held at a command potential \((V_C)\) of -75 to -80 mV, well below the activation range of \(I_A\) but sufficiently negative to steady-state deinactivate the current (Hayes et al., 2008). The initial \(V_C\) was held constant for all stimulus protocols through the duration of each experiment. The presence of \(I_A\) was confirmed by delivering increasing 400 ms step commands up to +90mV (relative to \(V_C\)). Neurons exhibiting a transient outward current that activated at subthreshold membrane potentials and inactivated above -40 mV after approximately 100-200 ms were considered to express \(I_A\) (Fig. 2. 2B).

Dendritic Ca\textsuperscript{2+} transients were then imaged during delivery of fast (150 ms duration) positive-going voltage ramps, starting at \(V_C\) and increasing to a final membrane potential capable of activating voltage-gated Ca\textsuperscript{2+} currents (mean = 7.8 ± 10.2 mV, \(n = 9\)). By using short-duration, quickly-increasing ramps we were able to elicit a supra-threshold Ca\textsuperscript{2+} response within the transient phase of
Iₐ activation (Fig. 2C). We then added 2 mM 4-AP to the bath perfusate and allowed 10 minutes for complete wash-in. Positive step commands as before were repeated to confirm blockade of Iₐ (Fig. 2B) and dendritic Ca²⁺ transients were imaged again while delivering the same voltage-ramps. The command stimulus was constant between control and 4-AP sweeps for each recorded neuron, but the peak amplitude achieved by the stimulus waveform (which is measurable in SEVC) nevertheless increased marginally across all neurons after addition of 4-AP because of increased effectiveness of the SEVC (type-1: 8.5 ± 2.2%, n = 5; type-2: 2.0 ± 1.5%, n = 4). Changes in peak amplitude of the stimulus waveform were significantly greater in type-1 neurons than those in type-2 neurons (n = 9, p = 0.0017). Input resistance in each cell was measured by taking the slope of the IV curve from the first 20 mV of increase above command potential—a range in which no apparent active conductances were elicited. Mean input resistance was 221 ± 97 MΩ in type-1 neurons and was not significantly different in type-2 neurons (181 ± 77 MΩ. n = 9, p = 0.52).

In order to visualize how voltage propagation through dendritic compartments is affected by a blockade of Iₐ, we measured the relative increase in Ca²⁺ indicator fluorescence (ΔF/F₀) evoked by voltage ramps before and after exposure to 4-AP (Fig. 2D). Measurements were sampled from proximal dendritic compartments (less than or equal to 33 µm from the soma) and distal dendritic compartments (greater than or equal to 69 µm from the soma) with a minimum distance between measurement sites of 54 µm. The mean proximal ROI distance from the soma was 14 ± 9 µm (n = 9) and the mean distal ROI distance from the soma
was 101 ± 21 µm (n = 9). The peak ΔF/F₀ values measured at each location were normalized to the amplitude of the control response in order to assess the relative change in fluorescence transient amplitude attributable to 4-AP effects (Fig. 2.2E). Among type-1 neurons, the amplitude of fluorescent Ca²⁺ transients elicited by voltage ramps increased significantly at proximal regions by 129 ± 102% (n=5, p=0.023) over the control amplitude and also increased significantly at distal regions by 282 ± 123% (n=5, p=0.003). The increase at distal regions in type-1 neurons was significantly greater than proximal regions (n = 10, p=0.032), suggesting a non-linear increase in the amplitude of calcium influx between perisomatic and distal dendritic compartments.

Among type-2 neurons, the amplitude of fluorescent Ca²⁺ transients did not change significantly at proximal regions (10 ± 22%, n = 4, p = 0.22) compared to control, while the amplitude at distal regions increased significantly by 37 ± 22% (n = 4, p = 0.022). The increase in the ramp-evoked Ca²⁺ transient at distal regions in type-2 neurons was not significantly greater than proximal regions (n = 8, p = 0.064) after 4-AP, which suggests that Iₜ does not play a significant role in governing dendritic depolarization of type-2 preBötC neurons, particularly compared to their counterparts the type-1 preBötC neurons (compare columns 1,2 to 3,4 in Fig. 2.2E).

These results demonstrate that a large increase in the voltage ramp-evoked Ca²⁺ transient occurs globally in type-1 neurons when 4-AP-sensitive currents are blocked (Fig. 2.2D, depicting raw ΔF/F₀ values above 100%), and that this increase is greatest at distal dendritic regions. The data imply either: 1) blocking
\(I_A\) diminishes the electronic decay of command voltages from the soma, which could be explained by removal of \(I_A\)-mediated shunting of the dendritic plasma membrane, or 2) blockade of \(I_A\) somehow increases recruitment of inward Ca\(^{2+}\) currents at distal dendritic sites. In either scenario articulated above, the data we present here cannot distinguish whether the observed changes in measured Ca\(^{2+}\) transient amplitude arise predominately from loss of somatic or dendritic \(I_A\). Thus, we sought to determine whether ionic membrane currents could be blocked on distal dendritic sites while minimally affecting the soma.

**Spatially-restricted application of channel blockers via iontophoresis**

To demonstrate that ion channels on selected sub-cellular regions of respiratory neurons in organotypic cultures can be pharmacologically manipulated, we conducted a positive control experiment in which Cd\(^{2+}\) was applied focally via iontophoresis to block Ca\(^{2+}\) channels in distal dendrites (Fig. 2.3). Simultaneous whole-cell patch-clamp and Ca\(^{2+}\)-imaging recordings were acquired from neurons in the preBötzC of organotypic slice cultures. Spontaneous network activity was depressed to prevent spurious signals by reducing extracellular K\(^+\) concentration from 8 to 2.5 mM and increasing extracellular Ca\(^{2+}\) concentration from 1.5 to 2 mM in the recording ACSF. Membrane potential was held between -55 and -60 mV to prevent spontaneous spiking. A pipette containing an aqueous solution of 200 mM CdSO\(_4\) was positioned with a robotic manipulator under visual control so that its tip aimed at distal dendritic regions, while maintaining a -5 nA holding current to prevent ion leakage.
Figure 2. Dendritic iontophoresis of Cd$^{2+}$ reduce dendritic Ca$^{2+}$-transients evoked by current pulses. A: 200 ms current pulses applied to a type-1 neuron evoking a spike train, and the resulting Ca$^{2+}$ transient in a dendritic compartment (red trace, ~130 µm from soma, at the Cd$^{2+}$ application site). Iontophoresis of Cd$^{2+}$ (2 min, 500 ms pulses, 1 Hz) reduced the dendritic Ca$^{2+}$ transient (black trace). B: Same neuron as in A, showing the spatial distribution of the Cd$^{2+}$ effect, expressed as attenuation (Red: 0 to Blue: 0.8). Note that the site of application (pipette insert) has the largest attenuation compared to more proximal sites along the dendrite. C: Group data (n=7) showing the attenuation of spike-train evoked Ca$^{2+}$ transients in response to Cd$^{2+}$ as a function of distance along the dendrite towards the soma. Black line is a linear fit.
Positive current through the patch-recording pipette was then injected at the soma in either square-wave pulses (400 ms, 0.1-0.3 nA, 1 Hz), causing repetitive spiking, or short repetitive current pulses (3 ms, 0.8-1.2 nA, 100 Hz; Fig. 2.3A) evoking trains of 10 to 25 action potentials. The stimulus was maintained between control and drug application in each cell. After acquiring control sweeps, continuous ejection pulses were delivered to the iontophoretic pipette (400-500 ms, 1 Hz, +5-10 nA). After allowing 30-60 seconds for equilibration of drug ejection, we repeated the imaging sweeps. Peak ΔF/F₀ values were sampled from two locations in each cell, separated by a minimum of 50 µm (n = 7 neurons): proximal and distal dendritic regions as before, as well as the soma. The decrease in Ca²⁺ transient amplitude after application of Cd²⁺ was normalized to control sweeps.

Distance between the tip of the iontophoresis pipette and center of each ROI was measured in the xy-plane. Since both the iontophoresis pipette and imaged cellular compartments occupy approximately the same focal plane, the difference in their positions estimated the distance in three-dimensional space between the point of drug application and measurement. Regression analysis revealed a linear relationship between the degree of Ca²⁺ signal attenuation and straight-line distance from the pipette (Fig. 2.3C, n = 14 measurements; y-intercept = 1.0, R² = 0.85; ANOVA F-value = 64.45, p < 0.001). This model suggests that less than 50% of the somatically evoked, dendritic Ca²⁺ transient is attenuated when the drug (Cd²⁺ in this case) is applied 79 µm away from the ROI. Although the mobility in the extracellular environment due to applied electric field undoubtedly
differs between Cd\textsuperscript{2+} and 4-AP, both drugs are extracellular ion channel blockers that are not taken up by cellular processes. Thus, to ensure that recorded somata remained unaffected by iontophoretic drug application, we maintained a minimum of 96 µm (and average of 121 ± 20 µm, n = 12 experiments) between the point of drug application and the nearest edge of somatic compartments in all subsequent local drug application experiments.

**Dendritic I\textsubscript{A} blockade increases Ca\textsuperscript{2+} response to somatically evoked stimuli in type-1 neurons**

To determine whether increases in the Ca\textsuperscript{2+} response of type-1 neurons due to 4-AP are mediated by dendritic I\textsubscript{A}, we applied 4-AP via iontophoresis to distal dendrite sites (Fig. 2.4, mean dendritic length from soma: 118 ± 21 µm, n = 12) in rhythmically active preB\text{"o}tC neurons and measured the amplitude of Ca\textsuperscript{2+} transients in response to trains of 10 action potentials triggered by somatic current injection. Whole-cell patch-clamp recordings of rhythmically active neurons in the preB\text{"o}tC of slice cultures were acquired and their membrane properties were tested in current clamp recording mode to test for the presence of I\textsubscript{A} and I\textsubscript{h}, and thus determine whether they were type-1 or type-2. At least 40 minutes after whole-cell break-in was allowed to pass for equilibration of dye diffusion at distal compartments. All imaging sweeps were performed in the presence of 2.5 mM K\textsuperscript{+} and 2 mM Ca\textsuperscript{2+}, which is the low-excitability ACSF as before.

We further added the ionotropic excitatory amino acid receptor antagonist NBQX (10-20 µM) to the perfusate to suppress excitatory network synaptic activity.
Figure 2. 4 Dendritic iontophoresis of 4-AP increases dendritic Ca\(^{2+}\) transients evoked by action potentials. A: A train of current pulses (10 pulses, 3 ms duration, 100 Hz) applied to a type-1 neuron evoking 10 spikes. The resulting Ca\(^{2+}\) transients in a dendritic compartment (~110 µm from soma, placed next to the 4-AP pipette) are shown in A, displaying evoked responses before and after (overlaid green traces) iontophoresis of 4-AP (7 min, 500 ms pulses, 1 Hz). Transients in the same neuron sampled from a dendritic site close to soma (blue traces) before and after 4-AP applied distally. B: Same neuron as in A, showing the color coded spatial distribution of the 4-AP effect, expressed as Ca\(^{2+}\) transient amplitude after 4-AP normalized to control (Blue: 105% to Red: 120%). Note that the site of application (pipette insert) has the largest increase in the Ca\(^{2+}\) transient compared to more proximal sites along the dendrite. C: Group data showing the Ca\(^{2+}\) transient amplitude after 4-AP application for dendritic sites that were either proximal or on other branches, and distal sites (i.e. in vicinity of the drug pipette) in both neuronal types (n=9 type-1 neurons, and n=3 type-2 neurons). Black dotted line is 100%, i.e. no change. Note that distal dendritic sites in type-1 neurons show a large increase in Ca\(^{2+}\)-transients after application of 4-AP.
In both control trials and local 4-AP application trials, ROIs were sampled from distal dendritic regions within 30 µm of the iontophoresis pipette. To verify that the effects of 4-AP observed near the site of iontophoresis was caused by local ion channel blockade, we also sampled the relative change in fluorescence from a presumably unaffected compartment (i.e. other dendritic branches, proximal sites on the branch of drug application, and the soma) located at least 100 µm away in the xy plane from the site of iontophoresis (mean distance: 139 ± 32 µm, n = 12). Pipettes containing 45 mM 4-AP dissolved in saline (165 mM NaCl, 0.2% Dextran-TMR, pH 7.5) were positioned at distal dendritic sites by visual guidance while maintaining a holding current of -5 nA to prevent drug leakage. During drug application trials, we applied continuous ejection pulses as before (400-500 ms, 1 Hz, +20-45 nA). In type-1 neurons, spike train-evoked Ca\(^{2+}\) transients at distal dendritic sites (near the point of drug application) increased substantially by 55 ± 29% (Fig. 2.4A-C, n = 9, p = 2.16E-4). Regions greater than 100 µm from the site of iontophoresis also increased in ΔF/ΔF₀ compared to the control response, which was also significant by statistical hypothesis testing but nonetheless represents a much less substantial change compared to the change measured in distal dendrites (7 ± 8%, n = 9, p = 0.017). However, the increase at sites near the point of iontophoresis was significantly greater than on other dendritic branches or the soma (Welch’s unequal variances t-test, n = 18, p = 4.4E-4).

Type-2 neurons did not show a consistent increase at the site of iontophoresis (Fig. 2.4C, 2 ± 10%, n = 3, p=0.34) or at sites >100 µm from the site of iontophoresis (5 ± 10%, n = 3, p=0.23). These results demonstrate that blockade
of $I_A$ on dendrites has a substantial effect on the size of distal dendritic $Ca^{2+}$ transients evoked by current pulses evoked at the soma in type-1 preBötC neurons, but that $I_A$ blockade has a negligible effect on dendritic $Ca^{2+}$ transients evoked at the soma in type-2 neurons.

2.3 DISCUSSION

A great deal is known about membrane properties of preBötC neurons, but almost all of that knowledge pertains to somatic ion channels and intrinsic membrane currents. We still lack a full understanding of how the active currents are distributed over the soma-dendritic membrane, including the dendrites where most of the synaptic input presumably arrives during respiratory network rhythmicity (Del Negro et al., 2011). Here we used organotypic slice cultures containing the preBötC to image $Ca^{2+}$-transients evoked by voltage increases propagating along extended dendritic lengths in rhythmically active neurons. The experiments show that an $I_A$ is present in the dendrites of type-1 neurons, which may influence synaptic integration.

The $I_A$ found in type-1 neurons of the preBötC likely influences the onset of the inspiratory burst phase during rhythmic network activity. Previous experiments show that bath applied 4-AP causes disordered inspiratory rhythms, and the $I_A$ has been measured at the soma in whole-cell and outside-out patch-clamp recordings (Hayes et al., 2008). As such, excitatory input arriving at the soma is to some degree inhibited by $I_A$ as long as the current is not steady-state inactivated. However, these previous measurements did not provide information
about the soma-dendritic distribution of $I_A$, which could influence how excitatory synaptic input is integrated both in the dendrites, and at the soma-axon hillock where spike initiation presumably occurs (Magee, 2000).

Interneurons in the preBötzC lack the laminar organization, planar dendritic arborization, or large-diameter dendrites that have made other cell types (e.g. pyramidal cells) amenable to methods of investigating active properties in dendrites (e.g. dendritic patch-clamp or $\text{Ca}^{2+}$ imaging). Organotypic slice cultures containing the preBötzC flatten and become more translucent than acutely prepared slices after 6 to 7 days in vitro, which improve optical qualities of the tissue and reduces the degree to which neuronal processes traverse the z-axis (Gähwiler et al., 1997; Phillips et al., 2016). Cultures are thus better suited than acute slice preparations for dendritic $\text{Ca}^{2+}$ imaging experiments. These slice cultures of the preBötzC retain bilateral rhythmicity, respond to known network rhythm modulators, and contain neurons whose behavior resembles that found in acute preparations which have been well-established as models of respiratory rhythm generation for over 25 years (Feldman et al., 2013; Funk and Greer, 2013; Phillips et al., 2016; Ramirez et al., 2004; Rekling et al., 1996; Smith et al., 1991). However, it remains uncertain whether or not the respiratory-like rhythm found in slice cultures containing the preBötzC arises from the same underlying cellular mechanisms as it does in acute slices that have been so widely exploited in studies of rhythm generation.

Membrane currents $I_A$ and $I_h$ can differentiate two distinct classes of rhythmically active neurons in the preBötzC, dubbed type-1 and type-2 (Rekling et al., 1996).
We found that these currents are well-preserved in culture and maintain a similarly segregated distribution. That is, the majority of rhythmically active neurons feature exclusively $I_A$ (i.e., type-1) or $I_h$ (i.e., type-2), and few express both $I_A$ and $I_h$ or neither of these two. To better understand how type-1 and type-2 neurons integrate synaptic input, we asked whether $I_A$ in rhythmically active neurons exists on dendrites and whether it has a significant impact on the ability for voltage to spread between somatic and distal dendritic compartments.

Blockade of $I_A$ in type-1 neurons results in a ~130% increase in the size of ramp-evoked $Ca^{2+}$ transients at proximal dendritic regions and a ~280% increase at distal dendritic regions. In contrast, type-2 neurons show no significant increase at proximal dendritic regions after bath application of 4-AP, while distal dendritic regions increased by ~40% over control. The global increase in the size of transients seen in type-1 neurons reflects either a change in the electrotonic compactness of the neuron—more of the somatically-triggered depolarization propagates from the soma to the distal dendrite—or that somatically triggered depolarization is less counteracted by $I_A$. Both interpretations are feasible because $I_A$ is an outward current whose polarity directly opposes depolarizing current longitudinally flowing from soma to dendrite and whose constituent ion channels in their open state may substantially change membrane impedance.

Bath application of 4-AP mediated a ~2-8% rise in the maximum amplitude of delivered voltage ramps in both type-1 and type-2 neurons, signifying a change in the strength of the SEVC. By some immeasurable degree, the consistent increase in ramp amplitudes definitely contributes to the observed change in
evoked transients on both proximal and distal dendrites. However, the error associated with type-1 neurons (i.e. ~8%) was significantly greater than that seen in type-2 neurons (i.e. ~2%), suggesting that blockage of \( I_A \) underlies a portion of the amplitude disparity. The strength of the SEVC, and by extension the ability to increase the membrane potential of the neuron, is thus correlated with the presence or absence of \( I_A \), supporting the hypothesis that it affects either electrotonic compactness or resists membrane depolarization.

The observed increase in ramp-evoked \( \text{Ca}^{2+} \) transients in type-1 neurons was also non-linear—distal sites increased ~2 fold more than proximal sites, suggesting that there is a disproportionate change in the amount of membrane depolarization occurring at distal versus proximal dendrites. Non-linear increases at distal dendritic locations could be explained as either an increase in the strength of space clamp mediated by the loss of \( I_A \) or the recruitment of active inward conductances at more distal dendritic sites (e.g. high-voltage activated \( \text{Ca}^{2+} \) channels). The possibility that active inward conductances may have been recruited at distal locations cannot be ruled out, particularly since type-2 neurons also displayed non-linear response increases. However, space clamp issues are exacerbated by the presence of voltage-dependent membrane conductances on distal compartments (Bar-Yehuda and Korngreen, 2008). An inability to fully compensate for \( I_A \) during SEVC voltage ramps in type-1 neurons indicates that these currents may be acting at a distance beyond the effective space clamp enforced near the soma. Ultimately, the explanation may be some combination of additional distally-located inward currents and an improved ability for SEVC to
clamp membrane potential further from the soma due to less inhibition of membrane depolarization caused by $I_A$, either somatically or also on dendrites.

The data from these experiments is unable to distinguish whether the apparent increase in electrotonic compactness, or reduced inhibition of membrane depolarization, is the result of solely somatic $I_A$ or additionally includes dendritic $I_A$. To definitively determine whether the increases in the amplitude of ramp-evoked $Ca^{2+}$ transients were caused by dendritic $I_A$ in conjunction with somatic $I_A$ we attempted to locally block $I_A$ on dendrites of rhythmically active neurons such that the soma would not be affected.

The dendrites of rhythmically active interneurons in the preBötC have a span of less than 300 µm (Picardo et al., 2013), and the maximum distance from the soma for which current injection evokes $Ca^{2+}$ transients was 153 µm. We employed iontophoretic drug ejection to ensure that we could restrict spread of ionic channel blockers applied at dendritic sites that were minimally 96 µm away from the soma. We tested our ability to focally block channels by using $Cd^{2+}$ as a positive control that would deliver complete knock-out of $Ca^{2+}$ transients at the site of application. Since our recordings were capable of capturing >100 µm of dendritic structures within the same focal plane, we took measurements at two cellular locations in each time series from a total of $n = 7$ neurons and pooled the data to generate a regression of signal attenuation against straight-line distance from the iontophoresis pipette (Fig. 2.3C). Although the ejection pipette was 96 µm away from the soma in one experiment, the average distance between the iontophoresis pipette and soma was $121 \pm 20$ µm, which implies that the ejected
drug could only be 36% effective at the soma compared to the dendritic ROI, according to our linear model (Fig. 2.3C). The mobility which 4-AP and Cd$^{2+}$ experience in an applied electric field differs, particularly since 4-AP does not carry a formal charge. While we were not able to precisely estimate the amount of 4-AP delivered to each cell, the control in Cd$^{2+}$ rather serves to demonstrate how consistent drug delivery can be maintained at ~100 µm distances from the soma trial-by-trial. It is indeed likely that some amount of 4-AP reached the soma in our recordings, but the results nevertheless strongly suggest that $I_{A}$ is present on dendrites where its effects are principally mediated.

Application of 4-AP to the dendrites of type-1 neurons resulted in a ~55% increase in spike train-evoked Ca$^{2+}$ fluorescence at the site of iontophoresis and ~7% increase in fluorescence in other compartments located either proximally on the same parent branch, at the soma, or on other dendritic branches occupying the same focal plane. The increase observed at other locations could indicate that the change in relative Ca$^{2+}$ fluorescence observed at the site of iontophoresis is in part explained by some degree of 4-AP reaching the soma. However, the integrity of action potentials generated at the soma served as a final control. Bath application of 4-AP causes the half-width at maximum of action-potentials to increase by approximately 80% (Hayes et al., 2008). Here, we excluded trials wherein action potentials became distorted after drug application (a sign of either 4-AP reaching the soma or cell instability). Thus, these results indeed suggest that the density of ionic channels giving rise to $I_{A}$
observed in type-1 neurons of the preBötC extends well beyond the soma into distal dendritic compartments.

I_A on the dendrites of rhythmically active preBötC neurons would be expected to counteract sparse excitatory synaptic events. It is important to distinguish that the enhancements in dendritic Ca^{2+} influx in the presence of 4-AP do not necessarily map one-to-one with a rise in voltage, but generally are indicative of changes in voltage, i.e., more Ca^{2+} influx indicates depolarization. The changes observed in ramp-evoked Ca^{2+} fluorescence were normalized to their control value in order to correct for confounds which could alter response linearity such as compartment volume, channel density, and the equilibrium of fluorescent indicator concentration (Yasuda et al., 2004). The non-linear increase in evoked Ca^{2+} transients at distal dendritic sites can be equally interpreted as a reduction in voltage decay between the soma and dendrites or as the recruitment of previously inhibited inward currents. Considering the overall global increase in Ca^{2+} response, I_A at minimum alters electrotonic compactness since all cellular compartments appear to charge in response to voltage stimuli more effectively. Propagation of voltage between sub-cellular compartments in type-1 neurons appears to be significantly affected by the availability of I_A whereas type-2 neurons have a more fixed relationship in the ability of transient somatic voltage increases to propagate through dendrites. Inactivation of I_A, which could be caused by sustained temporally-summated excitatory input, transitions type-1 neurons from a relative low-excitability state, in which excitatory synaptic input is presumably inhibited by outward current (i.e. I_A), to a high-excitability state that
appears to be significantly more electrotonically compact or resistant to membrane depolarization. This type of activity-dependent integration emphasizes why type-1 neurons expressing $I_A$ may be most critical in dictating the appropriately-timed onset of inspiratory burst cycles.

In conclusion, the subcellular distribution of $I_A$ in type-1 neurons extends to distal dendritic sites, and likely enforces a sublinear summation of input (i.e. EPSPs are inhibited) that is relieved by inactivation of $I_A$. This is evidenced by an apparent change in electrotonic compactness after blockade of $I_A$ by 4-AP. During rhythmic network activity, steady-state inactivation of dendritic $I_A$ could be achieved via building recurrent excitation during the pre-inspiratory phase of the inspiratory cycle. The presence of $I_A$ in the dendrites of type-1 neurons suggests that they are capable of limiting their excitability until network activity has grown during each respiratory cycle, and thus better suited for ordering the onset of inspiratory bursts.

2.4 METHODS

Ethical approval
The Department of Experimental Medicine at the Panum Institute approved all experiments and procedures according to protocols laid out by Danish Ministry of Justice and the Danish National Committee for Ethics in Animal Research.

Organotypic slice cultures
US Naval Medical Research Institute (NMRI) mice post-natal ages P3.5 to P6.5 days were anesthetized with isoflurane and immediately dissected in sterile-
filtered chilled artificial cerebrospinal fluid (ACSF) containing (in mM): 184 glycerol, 2.5 KCl, 1.2 NaH$_2$PO$_4$, 30 NaHCO$_3$, 5 HEPES acid, 15 HEPES base, 25 D-glucose, 5 sodium ascorbate, 2 thiourea, 3 sodium pyruvate, 10 MgSO$_4$, 0.5 CaCl$_2$; pH 7.3, equilibrated by bubbling with 95% O$_2$/5% CO$_2$. Transverse slices of the brainstem, 400 µm in thickness, were taken at the level of the preBötC using a vibrating microtome (Thermo Scientific Microm 650V, Waltham MA, USA). Anatomical markers, such as the principle loop of the inferior olive and obex of the fourth ventricle (Ruangkittisakul et al., 2011, 2014), were used to verify the rostral-caudal location of the preBötC and thus properly calibrate the slice. Cultures were prepared as previously described (Phillips et al., 2016) via the Stoppini interface method, placing transverse brainstem slices onto semi-porous culture well inserts (Millipore PIC03050, Darmstadt, Germany). Mounted preparations were maintained in sterile-filtered organotypic culture media containing: 50% Eagle’s MEM with Earle’s Salts, 25% Hank’s balanced salt solution, 25% horse serum, 2 mM GlutaMAX (Gibco), 200 U/mL penicillin, 5 µg/mL streptomycin, 25 mM HEPES and an additional 3.6 mM D-glucose. The osmolarity of the culture medium measured 320-340 mOsm with a pH of 7.25. The cultures were treated with 10 µM MK-801, an NMDA receptor antagonist, for the first three days in vitro to prevent ischemia-related cell death (Newell et al., 1990). Fresh culture media was supplied every 48 hours thereafter until experimentation. These slices were then kept in a sterile, humidified incubator at 35°C and atmospheric CO$_2$ concentrations during incubation.
**Electrophysiology**

Somatic whole-cell patch-clamp recordings were performed in current clamp and discontinuous single-electrode voltage clamp (dSEVC; sampling rate 1-2 kHz) using an AxoClamp 2B amplifier (Molecular Devices, Sunnyvale, CA, USA). Data were digitally acquired at a sampling rate of 10 kHz. Glass micropipettes were pulled from filamented capillary glass (O.D. 1.5 mm, I.D. 0.86 mm, Harvard Apparatus, Holliston, MA, USA) using a PUL-100 micropipette puller (World Precision Instruments, Sarasota, FL, USA) to a tip resistance of 4-6 MΩ. Patch pipettes were filled with a solution containing (in mM): 130 HCH₃SO₃, 130 KOH, 10 HEPES, 0.4 NaGTP, 4 Na₂ATP, 5 Na₂-phosphocreatine, 4 MgCl₂, 0.05 Alexa 594 hydrazide (Invitrogen, Carlsbad, CA, USA), 0.1 Fluo-8L (AAT Bioquest, Sunnyvale, CA, USA). The osmolarity of the patch pipette solution measured 310 mOsm with a pH of 7.3. Patch pipettes were visually guided to target neurons under visual control using ROE-200 micromanipulators (Sutter Instruments, Novato, CA, USA) on a fixed-stage upright microscope (modified Olympus BX51, Olympus Corporation, Tokyo, Japan) under 40x magnification.

Iontophoresis was performed using a IP-X5 instrument (Neuro Data Instruments Corp., New York, NY, USA). Glass micropipettes were pulled as described above to a tip resistance of 8-10 MΩ for single-drug trials. For experiments involving 4-aminopyridine (4-AP) pipettes were filled with an aqueous solution containing: 165 mM NaCl, 10 mM HEPES, and 0.2% tetramethylrhodamine-dextran. For inhibition of I₅ localized to dendrites, we included 45 mM 4-AP (pH 7.5). For experiments involving local iontophoretic application of Cd²⁺, an aqueous solution
containing exclusively 200 mM CdSO₄ was used. ACSF in Cd²⁺ experiments was free of phosphates to avoid precipitation. Pulses were delivered at 1 Hz (500 ms pulse duration) with the ejection currents: +20-30 nA to eject 4-AP, +5-10 nA to eject Cd²⁺.

During voltage-clamp recordings and dendritic glutamate iontophoresis, 1 μM TTX was added to the bath to block Na⁺ current. A-current was blocked with bath-applied 2 mM 4-AP when not applied locally. During spike train response experiments, 10 μM 2,3-Dioxo-6-nitro-1,2,3,4-tetrahydrobenzo[f]quinoxaline-7-sulfonamide (NBQX) disodium salt (Tocris Bioscience, Bristol, UK) was added to the bath to block Ca²⁺ transients arising from spontaneous excitatory synaptic transmission.

Rhythmic activity in slices was recorded in artificial cerebrospinal fluid (ACSF) containing (in mM): 124 NaCl, 3 KCl, 5 KH₂PO₄, 25 NaHCO₃, 25 D-Glucose, 1 ascorbic acid, 1 MgCl₂, 1.5 CaCl₂ with a pH of 7.4, equilibrated by bubbling with 95% O₂/5% CO₂. The final concentration of K⁺ was [K⁺]₀ = 8 mM, which globally elevates resting membrane potentials and increases the frequency of spontaneous respiratory rhythm. To identify Iₐ and I₉ in current clamp experiments, perfusion bath ACSF was exchanged after recording >15 minutes of rhythmic activity for a low excitability ACSF in which external K⁺ concentration was reduced from 8 to 3 mM and external Ca²⁺ was raised from 1.5 mM to 2 mM. These modifications reduced or stopped network rhythmic activity by globally hyperpolarizing neuronal membrane potentials and increasing the threshold for Na⁺ channel activation (Panaitescu et al., 2009). Low excitability ACSF contained
(in mM): 124 NaCl, 2.5 KCl, 1.25 NaH$_2$PO$_4$, 25 NaHCO$_3$, 25 D-glucose, 1 ascorbic acid, 1 MgCl$_2$, 2 CaCl$_2$, with a pH of 7.4, equilibrated by bubbling with 95% O$_2$/5% CO$_2$.

**Ca$^{2+}$ imaging**

The region of the organotypic culture (see Fig. 2.1 in Phillips et al. 2016) containing the preBötC was targeted via anatomical landmarks and visualization of synchronized rhythmic network activity during whole-slice Ca$^{2+}$ imaging. The membrane-permeable fluorescent Ca$^{2+}$ indicator, Fura-2 AM (AAT Bioquest), was bath loaded prior to each experiment. Culture inserts were submerged in 1.5 mL of loading solution containing 35 µM of a 1 mM stock solution of Fura-2 AM in DMSO along with 0.05% pluronic acid dissolved in low excitability ACSF. Slices were incubated in loading solution at room temperature for 30-40 minutes then moved to the recording chamber and allowed 15 minutes of recovery and additional de-esterification prior to imaging and patching. Preheated and oxygenated ACSF was perfused into the recording chamber during recording at 2 ml/min and 29° C.

Fluorescent Ca$^{2+}$ activity was recorded in wide-field on a fixed-stage upright microscope (modified Olympus BX51), illuminated by a metal halide light source PhotoFluor II (89North, Burlington, USA) or a LED light source, M470L2 (Thorlabs, Newton, NJ, USA). Red and green channel fluorescence was visualized using a dual-bandpass filter set (Chroma 59022: excitation dual band-pass 450-490 nm/555-590 nm, emission dual band-pass 500-543 nm/603-665 nm). Red and green channels were separated during acquisition by manually
exchanging an additional excitation filter in the light train (Semrock FF01: band-pass 565-605 nm; Semrock FF02: band-pass 457-487 nm, Rochester, NY, USA). Time series acquisition was performed with a sCMOS camera (Neo DC-152Q, Andor Tehcnology, Belfast, UK) controlled by SOLIS software (Andor Tehcnology). Imaging protocols employed 10x (NA 0.3) and 40x (NA 0.8) water immersion objectives. Time series were acquired at 10-50 Hz. At least 30 minutes of perfusion was allowed after break-in during whole-cell patch clamp to allow equilibration of intracellular Ca\(^{2+}\) dye. Dark current—the background sensor noise in the absence of illumination—was sampled prior to the start of each acquisition and subtracted from all imaging sweeps.

**Data analysis**

Dendritic Ca\(^{2+}\) transients were calculated as the percent change in fluorescence relative to baseline values (\(\Delta F/F_0\)). Background was subtracted frame-by-frame, taken as the mean background fluorescence immediately adjacent to the dendritic ROI. Dendritic ROIs were hand drawn along neural processes, extending for no more than 10 µm in length. The mean pre-stimulus fluorescence in the dendritic ROI was taken as \(F_0\). For each condition, 4-6 sweeps were acquired with 10-30 second intervals, and subsequently averaged after calculation of \(\Delta F/F_0\).

Electrophysiological data were acquired using pClamp 10.0 (Molecular Devices, Sunnyvale, CA, USA) and subsequently analyzed using custom scripts written in Igor Pro 6 (Wavemetrics, Tigard, OR, USA).
Unless otherwise stated, statistical values are given as mean ± SD. Student’s t-test was used for statistical comparisons of one- and two-sample populations. In cases where equal variance between samples could not be assumed, Welch’s correction was applied and is indicated. Fisher’s exact test was used to assess the distribution of neuronal membrane currents. All statistical tests were performed using Origin 9.1 (OriginLab Corp., Northampton, MA, USA).
2.5 REFERENCES


CONCLUSIONS

Breathing is vital to life, obviously, and understanding its neural origins is an important problem for physiology and neuroscience. Respiration is controlled by neuronal populations distributed throughout the brainstem, of which the preBötzinger complex is predominant as the source of the inspiratory rhythm, coordinating all other respiratory phases (e.g., post-inspiration and expiration) as well as most orofacial behaviors, e.g., whisking, sniffing, licking, chewing, and swallowing (Kleinfeld et al., 2014; Moore et al., 2013, 2014). To understand how breathing rhythm is generated, the brainstem has been reduced to isolate the most essential network circuitry necessary for rhythmogenesis (Funk and Greer, 2013; Rekling and Feldman, 1998; Smith et al., 1991; Suzue, 1984). This reductionist approach culminated in the development of acute medullary slice preparations, which isolate just the preBötzinger complex and sufficient premotor and motor neurons to generate rhythm and measurable inspiratory related motor output (Smith et al., 1991). These slices for over 25 years have substantially improved the ability of investigators to investigate the cellular and synaptic properties of respiratory interneurons, and have also been used profitably in molecular genetic analyses of these central circuits (Bouvier et al., 2010; Del Negro et al., 2002; Feldman et al., 2013; Gray et al., 1999, 2010; Rekling et al., 1996; Tan et al., 2008). However, acute medullary slices come with several caveats which have
restricted experimental design—namely, slices are at most viable for less than one day (Funk and Greer, 2013), and morphological characteristics of preBötzinger complex interneurons complicate studying synaptic integration in dendrites. The methodologies developed herein and experiments presented in this dissertation have attempted to ameliorate the drawbacks inherent in acute slice studies in order to both expand possible experimental approaches and to evaluate the role of a particular membrane conductance ($I_A$) as it relates to a fundamental—but incompletely understood—component of inspiratory rhythm generation: integration of excitatory synaptic activity into inspiratory bursts occurs primarily in dendrites (Pace and Del Negro, 2008; Pace et al., 2007; Rekling et al., 1996).

Chapter 1 presented a novel organotypic slice culture containing the preBötzinger complex and demonstrates that its behavior closely resembles analogous acute slices. The rhythm is generated by surviving interneurons whose electrophysiological behavior, synaptic projections and frequency-modulating receptor types (i.e. neurokinin-1 receptors, $\mu$-opioid receptors) remain intact. Calcium imaging from these cultured preparations highlights improved optical qualities, allowing isolation of somatic calcium transients from up to 200 neurons at a time under 10x magnification and subcellular measurements of fluorescence intensity under 63x magnification (see Figure 1.3).

Organotypic slice cultures containing the preBötzinger complex are a useful alternative to acute slice preparations with the added advantage of 1 to 4 weeks of viability and improved optical qualities for imaging. By increasing the window of time in which experiments may be conducted, investigators can utilize
molecular techniques that require multi-day incubation periods (e.g. viral transduction of recombinant DNA or non-viral transfection of plasmid DNA). The ability to express genes of interest or exogenous DNA (e.g. coding for fluorescent proteins) and subsequently record from rhythmically active preparations is particularly useful to respiratory neurobiologists, permitting rapid identification of cell types of interest and the use of genetically encoded molecular tools (e.g. optogenetics). It is indeed possible to stereotaxically inject viral vectors carrying recombinant DNA (e.g., adenovirus) into the preBötC of living animals. However, such injections are typically performed in juvenile or adult mice and then assayed approximately one week later in vivo or histologically. Although cellular recordings of rhythmic activity have previously been achievable in acute slice preparations taken from juvenile mice, thicker tissue sectioning (e.g. ~700 μm) is required to capture the preBötC and hypoglossal motor nuclei in a single slice and such preparations do not reliably produce motor output beyond P21, making them more difficult to study (Funk et al., 1994; Ramirez et al., 1996). To the best of my knowledge, an adult slice that retains the preBötC and remains viable and rhythmically active in vitro, has not been accomplished. As such, the usefulness of acute slice preparations lies between embryonic stages up to P14 (realistically this window is limited to P4, and P14 is not nearly as advantageous for recording and imaging), which is well before targeted in vivo injections can affect respiratory circuits via transduction of recombinant DNA. Alternatively, transgenic animals can be used to express exogenous DNA or to modify genes of interest or express certain reporter proteins, but transgenic models require development of
at least one (and often two) viable mutant mouse strains for each experimental objective. Further, promoter-driven expression or knock-out occurs genome-wide, which modifies the entire brain and CNS experimental confounds. Thus, the options for vector DNA expression are limited for *in vitro* experiments. Organotypic slice cultures, on the other hand, do not suffer from these restrictions. Viral transduction has already been demonstrated in rhythmically active slice cultures containing the preBötC (Forsberg et al., 2016)(Fig. C.1; Rekling unpublished). Transfection in organotypic cultures also need not be viral-based nor ubiquitous. Plasmid DNA can be delivered non-virally (Murphy and Messer, 2001), in a region specific manner (Arsenault et al., 2014; Wickersham et al., 2007), or even to single cells via electroporation (Nguyen et al., 2012; Rathenberg et al., 2003). The usefulness of organotypic slice cultures containing the preBötC has likely only begun to be realized.

![Image](example.png)

Figure C.1 Adeno-associated viral transduction of red fluorescent protein in slice cultures containing the preBötC. Left panel: Cropped image focusing on strong expression of red fluorescent protein in a single neuron taken at 40x magnification. Note expression throughout the soma-dendritic morphology. Right panel: View of the same neuron uncropped. Note widespread expression of the red fluorescent protein in multiple cells.
Chapter 2 demonstrated that the subcellular distribution of a transient outward K$^+$ current, $I_A$, extends onto the dendrites of type-1 rhythmically active preBötz neurons. Dendritic $I_A$ may inhibit excitatory input that occurs sparsely in rhythmic neurons, which would act to suppress these inputs (as long as $I_A$ remains deinactivated). That scenario may characterize the interval between inspiratory bursts. Nonetheless, as activity increases among preBötzC neurons, the temporal summation of repetitive input will cause steady-state inactivation of $I_A$, which would facilitate these inputs and promote synchronous burst generation in the network. Excitatory activity occurring when $I_A$ steady-state inactivates experiences a lesser degree of current inhibition when propagating from dendrites to the site of action potential initiation. Type-1 neurons thus appear to be uniquely equipped to promote recurrent excitation and thus periodic burst output, while inhibiting spontaneous excitatory input during interburst intervals.

Over 20 years ago, dendrites were predicted to contain intrinsic membrane conductances necessary for amplification of synaptic input in rhythmically neurons of the preBötzC. Nearly 10 years ago, amplification of synaptic input was directly linked to metabotropic glutamate receptors in dendrites (Pace and Del Negro, 2008; Pace et al., 2007). Synaptic integration occurring on dendrites has since been observed and measured in the context of rhythmic activity (Del Negro et al., 2011). However, the subcellular distribution of currents in rhythmically neurons has until now been undefined. The manner in which specific ionic membrane currents in dendrites (e.g., $I_A$) might interact with excitatory events known to also occur in dendrites (e.g. drive amplification) has similarly not been
tested. While the experiments performed here speculate upon the likely interactions occurring in dendrites due to $I_A$ (i.e. excitatory inhibition), the effects of $I_A$ on excitatory input in other systems have been documented (Hoffman et al., 1997; Magee, 2000; Magee et al., 1998). For instance, dendritic $I_A$ in Purkinje neurons is modulated via group I mGluRs and inhibits calcium spiking by high-voltage-activated calcium channels. These calcium spikes evoke bursts of action potentials recorded at the soma (Otsu et al., 2014). Although amplifying currents on dendrites in rhythmically preBötC neurons appear to be activated by a non-specific cation current ($I_{CAN}$), group I mGluRs do indeed promote drive amplification (Pace and Del Negro, 2008), and co-localize with $I_A$ as demonstrated here. Experiments involving repetitive excitation (i.e. trains of single EPSPs) or dendritic glutamate application, both paired with blockade of $I_A$ localized to dendrites, could confirm that $I_A$, in its deinactivated state, can diminish the effects of sparse excitatory inputs. Additional pharmacology could likewise test whether mGluRs interact with $I_A$.

Broadly speaking, the experiments in Chapter 2 highlight the complexity of integrative processes occurring in rhythmically active neurons. Moreover, this may not be exclusive to type-1 neurons. The hyperpolarization-activated inward current ($I_h$) is prominently featured in type-2 neurons (Picardo et al., 2013; Rekling et al., 1996), is known to be expressed on dendrites (Lörincz et al., 2002; Notomi and Shigemoto, 2004), and could likewise inhibit temporal summation of excitatory input (Magee, 1998; Stuart and Spruston, 1998; Williams and Stuart, 2000).
The organotypic culture model of inspiratory rhythm generation presented here recapitulates the behavior of acute *in vitro* preparations, provides a means by which experimenters can observe integrative processes occurring on dendrites and additionally lengthens the time-scale on which experiments can be conducted. This in turn permits future use of genetically encoded molecular tools such as protein indicators (e.g. calcium or voltage sensors) and light-activated ion channels (e.g. channelrhodopsins) that can be expressed in a cell- or region-specific manner. As proof of its utility, the culture model here has permitted investigation of integrative properties in dendrites of type-1 neurons, first predicted over 20 years ago. Until now, these properties were evidenced to exist, but only by indirect means or methods that could not elaborate in detail on the interactions of underlying active membrane currents. Here we show how I_A can influence the behavior of type-1 neurons through inhibition of membrane depolarization on their dendrites, which helps to reinforce their putative role as reliable rhythm initiators and provides valuable insight about the neural control of respiratory rhythm.
REFERENCES


