Characterization of Developmental Changes in Spontaneous Electrical Activity of Medial Superior Olivary Neurons Before Hearing Onset With a Combination of Injectable and Volatile Anesthesia

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In this work the impact of two widely used anesthetics on the electrical activity of auditory brainstem neurons was studied during postnatal development. Spontaneous electrical activity in neonate rats of either sex was analyzed through a ventral craniotomy in mechanically ventilated pups to carry out patch clamp and multi-electrode electrophysiology recordings in the medial region of the superior olivary complex (SOC) between birth (postnatal day 0, P0) and P12. Recordings were obtained in pups anesthetized with the injectable mix of ketamine/xylazine (K/X mix), with the volatile anesthetic isoflurane (ISO), or in pups anesthetized with K/X mix that were also exposed to ISO. The results of patch clamp recordings demonstrate for the first time that olivary and perilivary neurons in the medial region of the SOC fire bursts of action potentials. The results of multielectrode recordings suggest that the firing pattern of single units recorded in K/X mix is similar to that recorded in ISO anesthetized rat pups. Taken together, the results of this study provide a framework to use injectable and volatile anesthetics for future studies to obtain functional information on the activity of medial superior olivary neurons in vivo.

Keywords: ketamine, isoflurane, MNTB, in vivo electrophysiology, multi-unit activity

INTRODUCTION

The accurate organization of neuronal circuits is established during development through activity-dependent and activity-independent processes that involve the reorganization and fine-tuning of immature synaptic and cellular networks (Goodman and Shatz, 1993; Hanson and Landmesser, 2004; Kirkby et al., 2013). Before the onset of sensation, spontaneously active cells and signaling mechanisms in sensory organs have been identified as drivers of bursts of neuronal activity that are implicated in activity-dependent refinement of auditory and visual systems in different...
vertebrate species (Maffei and Galli-Resta, 1990; Kotak and Sanes, 1995; Lippe, 1995; Jones et al., 2007; Tritsch et al., 2007; Sonntag et al., 2009; Seabrook et al., 2017; Gribizis et al., 2019). In the auditory system of altricial rodents, calcium action potentials (APs) in cochlear inner hair cells (IHCs) initiate minibursts of APs in auditory neurons before the onset of hearing (Tritsch et al., 2007, 2010; Wang et al., 2015). During the prehearing period, IHCs are transiently innervated by direct axo-somatic efferent synaptic contacts from medial olivocochlear (MOC) neurons located in the brainstem superior olivary complex (SOC; Warr and Guinan, 1979; Simmons et al., 1996). MOC innervation is cholinergic (Glowatzki and Fuchs, 2000; Katz et al., 2004; Gómez-Casati et al., 2005) and mediated by postsynaptic acetylcholine nicotinic receptors (nAChR) containing α9 and α10 subunits (Elgoyhen et al., 1994, 2001; Weisstaub et al., 2002; Lipovsek et al., 2012). The nAChR is coupled to the activation of small-conductance calcium-activated SK2 potassium channels expressed in the IHCs, which mediate the hyperpolarization of IHC membrane potential in response to MOC efferent activation (Glowatzki and Fuchs, 2000). Thus, it has been proposed that MOC efferent-mediated inhibition might contribute to pattern trains of IHC calcium APs during the critical developmental period preceding hearing onset (Kros et al., 1998; Glowatzki and Fuchs, 2000; Marcotti et al., 2003; Johnson et al., 2011; Sendin et al., 2014; Moglie et al., 2018).

In recent years, different lines of genetically manipulated mouse models were used to study how modulation of the cochlear pacemaker affects the maturation of central auditory neurons and synapses. In the brainstem medial nucleus of the trapezoid body (MNTB), electrophysiological techniques were employed in slices and in vivo recordings of anesthetized α9 nAChR mutant mouse pups (Clause et al., 2014; Di Guilmi et al., 2019). Despite this progress, we identified a major discrepancy in published studies based on the use of different types of anesthetics in different rodent species. The two most widely used anesthetics in brainstem studies are the injectable mix of ketamine/xylazine (K/X mix) in mice (Sonntag et al., 2009; Clause et al., 2014; Di Guilmi et al., 2019), and the volatile anesthetic isoflurane (ISO) in rats (Tritsch et al., 2010; Crins et al., 2011; Sierksma and Borst, 2017). Although different factors are taken into account to choose the type of anesthetic, few studies have addressed its potential effects on in vivo spontaneous neuronal activity. For example, although previous studies have identified burst firing units in mice and rats (Sonntag et al., 2009; Tritsch et al., 2010; Clause et al., 2014), only one study in ISO-anesthetized rats reported regular firing units (Tritsch et al., 2010). This raises the possibility that the firing pattern of MNTB and other auditory brainstem neurons could be affected by the type of anesthetic used. Recent works in awake animals do not settle this issue, because optical calcium reporter fluorescence techniques were employed in auditory midbrain and cortex (Babola et al., 2018, 2020; Wang H.-Y. et al., 2020), and when electrophysiology experiments were used, they were performed in the somatosensory cortex of awake mice (Wang Y. et al., 2020).

In this work, we studied neuronal activity with electrophysiology techniques for single unit recording and ensemble multi-unit recording in the SOC of neonate rats. We used K/X mix to induce and maintain rat pups in a stable anesthetized state, while the percent ISO was titrated with precision to study its inhibitory effects on single unit and multiunit activity of medial superior olivey neurons. We also used the two anesthetics independently to monitor the overall activity of a population of neurons in animals at different postnatal ages, and ISO to identify the location of neurons with different firing properties. We found that burst and regular firing units are present in K/X mix anesthetized rat pups, that when ISO is added to K/X mix anesthetized pups it has inhibitory effects on the two types of units, and that some regular firing cells seem to be resistant to the inhibitory effects of maximal doses of ISO. Our results also demonstrate that despite the suppressive effects of anesthetics, the ensemble electrical activity of superior olivary neurons recorded in K/X mix or in ISO-anesthetized rat pups ramps up during development.

MATERIALS AND METHODS

Animal Housing and Breeding

The Institutional Animal Care and Use Committee of the City College of New York specifically reviewed and approved this study. The cohorts of adult Wistar rats used in this study were obtained from a commercial supplier at postnatal age 65 (P65, Charles River). Breeding trios of one male and two females were set for 5 days. At the completion of the breeding period, males were removed from the study while females were housed in pairs for 14 more days, and then individually until they gave birth. Pups were used from the day of birth (P0) to 12 days after birth (P12). Experiments were designed to minimize the number of animals used.

Ventral Surgery and Brain Processing

A total of 85 pups from 37 litters were prepared for surgery to expose the ventral skull for electrophysiology experiments. The surgical procedures used are those described by Rodriguez-Contreras et al. (2008), with minor modifications. Neonate rat pups were initially anesthetized by an intraperitoneal injection of a mixture of ketamine hydrochloride (0.1 mg/g body weight; Ketathesia) and xylazine hydrochloride (0.005 mg/g body weight; Anased), here referred as K/X mix. After anesthesia induction, animals were tracheotomized, intubated, and mechanically ventilated using a MiniVent type 845 ventilator (Harvard Apparatus). During surgery, anesthesia was carefully monitored on the basis of pedal reflexes, and maintained by supplementary injections of one-third of the initial dose of K/X mix as necessary (Sonntag et al., 2009). For the subset of experiments described in Figure 2, animals were initially anesthetized inside an induction chamber with 3% isoflurane (ISO) carried in oxygen, and maintained anesthetized with 1.5% ISO carried in oxygen, delivered via a nose cone or a direct connection to the mechanical ventilator. A small ventral craniotomy (1.5 × 1.5 mm) was performed on each pup, and the brain vascular landscape constituted by the basilar artery and the anterior inferior cerebellar artery was exposed. The dura was carefully removed prior to recording. Body temperature
was controlled during surgery and recording with a heating pad set to 37° (FHC, Maine, United States). After successful recordings were obtained, animals received an overdose injection of Euthanol, and were removed from the setup to be perfused with a solution of 4% paraformaldehyde in 0.1 M phosphate buffer. Brains were removed from the skull and processed for histology to be sectioned in 60 μm thick slices. Brainstem slices were mounted onto glass slides and counterstained with DAPI prior to applying mounting medium and a glass coverslip. Brain sections were imaged with an LSM800 confocal microscope located at the CCNY imaging core facility.

**Patch Clamp Recordings**

Recordings were made at a depth of 200–600 μm from the pial surface using borosilicate glass pipettes (5–8 MΩ) filled with artificial cerebrospinal fluid (ACSF) containing (in mM): 125 NaCl, 5 KCl, 2 MgSO₄, 2 CaCl₂, 10 D-glucose, 10 HEPES, adjusted to pH 7.4 with NaOH, or intracellular solution containing (in mM): 126 K-glutamate, 20 KCl, 10 Na₂₃-phosphocreatine, 4 MgATP, 0.3 Na₂GTP, 0.5 EGTA, and 10 HEPES (310 mmol.kg⁻¹), adjusted to pH 7.2 with KOH. Extracellular potentials were detected in the loose-patch configuration, while intracellular potentials were recorded in the whole-cell configuration. Pipettes were advanced from the brain surface in large steps of 50 μm to a depth of 200 μm using high positive pressure (300 mbar). When in cell search mode, pressure was reduced to 30 mbar and the pipette was moved in 1.5 or 2 μm steps. The pipette tip resistance was monitored with a 10 mV test pulse in voltage clamp or when a shift in the DC was noted in current clamp. When the tip resistance increased suddenly, suggesting that a putative cell was encountered, pressure was released and slight negative pressure of 30–50 mbar was applied until a low resistance seal was formed (10–50 MΩ) for loose-patch recordings, or until a high resistance seal was formed (1–10 GΩ) for whole-cell recordings. High-resistance seals were broken by applying fast bouts of negative pressure. Whole-cell recordings were not corrected for the liquid junction potential. Voltage was recorded with pClamp9 software using an Axopatch 200B amplifier, filtered at 2–5 kHz and sampled at 20 kHz with a 90 s period in 1.5% ISO and two washout bins. The effect of ISO was evaluated in multi-electrode recordings by using a procedure similar to the first protocol described above, except that between the baseline recording and the first two blocks in 1.5% ISO, a 90 s period in 1.5% ISO was added.

**Data and Statistical Analyses**

Unless indicated, data is presented as mean ± sem. In all cases, n indicates the number of units or cells tested, with the exception of Figure 3 where n indicates the number of animals. Extracellular single-unit data was filtered in two steps. First, to eliminate 60 cycle electrical interference (0.6 Hz, 3 dB bandwidth); and second, with a high-pass Bessel (8-pole) filter with a −3 dB cutoff of 5 Hz using Clampfit 10.6 (Molecular Devices). All data was analyzed using NeuroMatic in Igor Pro software (WaveMetrics; Rothman and Silver, 2018). APs were detected using a positive threshold set to six times the standard deviation of the baseline noise. Single-unit recordings where an obvious gradual decrease or rundown in spike amplitude occurred were excluded from analysis. Multielectrode multi-unit data was exported as NEX files for analysis in Plexon software (Offline sorter 4.4 and Neuroexplorer). APs were detected using a negative threshold set to six times the standard deviation of the baseline noise. Multiunit activity (MUA) collected in every channel was considered. All statistical tests were performed with Statistica 7.0 software (Stat Soft; RRID:SCR_014213). Non-parametric Kruskall-Wallis (K-W) test was used. Values of p < 0.05 were considered significant.

**Drugs and Reagents**

All drugs and reagents were purchased from Sigma-Aldrich (RRID:SCR_008988).

**RESULTS**

**Isoflurane Inhibits Burst and Regular Firing Units in the Ventral Brainstem of K/X Mix-Anesthetized Rat Pups**

Single-unit recordings were performed in five K/X mix-anesthetized rat pups before hearing onset, and the effects of 1.5% isoflurane were evaluated on the firing properties of six single-units (one unit from a P3 pup and five units from four P6 pups). Two of the units showed burst firing...
and four units showed regular firing patterns under baseline conditions (exemplar recordings are shown in Figures 1A–C). These recorded units had variable responses to ISO. The activity of the two burst firing units was inhibited to 96 and 99.6% compared to baseline and did not recover during washout. The activity of three out of the four regular firing units was partially inhibited to 34 ± 21% compared to baseline, and only in one unit recovered to baseline levels during washout. Lastly, one of the four regular firing units slightly increased its activity by 2% in ISO compared to baseline, but its activity was suppressed to 19% during washout. These inhibitory and potentiating effects of ISO on single-unit firing properties were observed in the inter-spike-interval histograms shown in Figures 1D–F. Small changes in the waveform of recorded units were also noted (insets in Figures 1D–F).

To address the variation of 1.5% ISO effects on recorded units, additional experiments were performed on thirteen single-units recorded from two P3 pups, two P6 pups, and three P9 pups by increasing the ISO dose from 0 to 4.5% in 1.5% steps. It was found that two P3 units, two P6 units and four P9 units showed clear signs of dose-dependent inhibition at 4.5% ISO compared to baseline. These cells did not recover during the washout period (Figure 1G). In a total of five additional single-units from pups of all ages, evidence of potentiation or resistance to 4.5% ISO was found. In three of these units, a small inhibitory effect was observed during the washout period (Figure 1H). Three of these

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**FIGURE 1** | ISO inhibit single unit activity in K/X mix anesthetized pups. (A) Burst firing unit from a P6 rat. (B) Regular firing unit from a P3 rat. (C) Regular firing unit from a P6 rat. (D) Effect of ISO on the inter-spike interval (ISI) distribution of single unit shown in (A). (E) Effect of ISO on ISI distribution of single unit shown in (B). (F) Effect of isoflurane on ISI distribution of single unit shown in (C). (G) Inhibitory effect of ISO on single unit activity recorded at P3 (green), P6 (blue) and P9 (magenta). (H) ISO resistant single units recorded at P3, P6 and P9. (I) Exposure to 5% ISO inhibited 3 of 5 single units shown in (H). ISI = inter-spike interval. Asterisk in D indicates the pre-spike in this complex waveform. Inset waveforms in (D) are averages of 269 and 11 action potentials in baseline and isoflurane, respectively. Inset waveforms in (E) are averages of 36, 11, and 302 action potentials in baseline, isoflurane and washout, respectively. Inset waveforms in (F) are averages of 2,538, 2,596, and 2,046 action potentials in baseline, isoflurane and washout, respectively. Lines and symbols in (H,I) identify the same single units under different isoflurane conditions. Vertical dashed lines in (G–I) indicate the onset and offset of isoflurane application.
it is not recover during the washout period. The remaining two units from P6 and P9 pups showed signs of resistance and potentiation even at such high dosage (Figure 1I).

Altogether, these experiments show that burst and regular firing cells are present in K/X mix-anesthetized rat pups, that most burst and regular firing units are inhibited permanently by ISO, while some regular firing units appear to be resistant to inhibition and show indications of potentiation after ISO is delivered to K/X mix-anesthetized rat pups. Resistance and potentiation to ISO was observed more frequently in P6 and P9 rat pups.

Heterogeneous Population of Single-Units: Inactive Versus Active Neurons

It was noted that from a total of nineteen recorded single-units, five units fired in a burst type pattern, while the rest fired in a regular spike pattern. Furthermore, three of the burst firing units did not show complex waveforms adjudicated to MNTB neurons. These observations raised the possibility that other SOC neurons may fire burst type or regular type patterns. One caveat of loose-seal patch clamp recordings is that cells cannot be identified, unless iontophoretic methods are used to label them (Pinault, 1996; Cid and de la Prida, 2019). As an alternative to identify the morphology and location of recorded cells, whole-cell current clamp recordings were performed in forty ISO anesthetized pups between ages P0 and P11 (Figure 2). Active and inactive cells were identified in current clamp recordings. Active cells fired APs in burst ($n = 10$) or regular type ($n = 5$) patterns, while inactive cells did not fire APs at rest ($n = 25$; Figure 2A). Two kinds of inactive cells were further identified based on their response to depolarizing current injection. Some cells fired APs ($n = 13$), while other cells did not ($n = 12$; Figure 2B). Therefore, inactive cells were classified as able or unable to fire APs (Figure 2C). After histological processing, the morphology and location of twenty-three recorded cells was confirmed in the medial region of the SOC (Figure 2D). Eight burst firing cells were observed within and outside the boundaries of SOC nuclei such as the MNTB and the MSO/SPN; two regular spiking cells were observed in the dorsal region outside the boundary of the MNTB; two inactive cells were observed within the boundaries of the MNTB; and eleven inactive cells were found at ventral, medial, lateral and dorsal locations outside the boundaries of the MNTB and MSO/SPN.

In sum, these experiments show that ISO-anesthetized rat pups have active and inactive neurons in the SOC. Several cells located in the medial region of the SOC fire APs in burst or regular patterns. However, while burst-firing cells were observed within and outside the boundaries of SOC nuclei, regular firing cells were observed only outside the boundaries of SOC nuclei. Some inactive cells are capable of firing action potentials upon depolarization of their membrane potential, while others cannot fire action potentials suggesting irreversible inhibition or a different cellular phenotype (i.e., non-neuronal glial).

Multiunit Activity Increases During Postnatal Development

Next, the effect of the injectable and volatile anesthetics on the ensemble activity of the medial SOC was evaluated separately
with multi-electrode probes in pups aged between P0 and P12 \( (n = 29); \text{Figure 3A}. \) Despite the large variability of spontaneous MUA within each developmental group, the overall MNTB spontaneous activity ramped up during development under K/X mix \( (n = 14); \text{Figures 3B,C, left panel} \) or ISO \( (n = 15); \text{Figures 3B,C right panel} \). It is important to note that the mean firing rate was similar under both conditions. Statistical analysis showed only differences under ISO displaying lower activity at P0 compared to P9 \( (K-W, p = 0.04; \text{multiple comparisons, } p < 0.05; n = 3) \). Taken together, these results suggest that the overall level of spontaneous electrical activity increases during postnatal development and that the maximal ensemble firing rate was similar between the two anesthetics.

**Developmental Profile of ISO Inhibition**

To examine whether ISO may act over the anesthetic effect of K/X mix and considering that both anesthetics have different synaptic targets \( (\text{MacDonald et al., 1987}; \text{Wang Y. et al., 2020}) \), we implemented the recording protocol illustrated in Figure 4A. Figure 4B displays three examples of MUA at different developmental ages (P3, P6, and P9) along the experimental protocol. In all cases, application of 1.5% ISO depressed the spiking activity, and a partial or total washout could be observed. The activity level was quantified after application of 1.5% ISO to K/X mix baseline \( (\text{Figure 4C}) \). P0 was excluded from the statistical analysis due to the activity being too small and the estimation of inhibition may not be reliable. Statistical analysis showed that ISO at 10 min had a greater initial inhibition at P6 in comparison with P12 \( (K-W, p = 0.007; \text{multiple comparisons, } p < 0.05; n = 3) \). In conjunction with loose-seal patch clamp recordings \( (\text{Figure 1}) \), these experiments show that the inhibitory effect of ISO is heterogeneous before hearing onset.

**DISCUSSION**

General anesthetics are widely used for surgical procedures in animal models and sometimes are required during in vivo experiments. For example, recent studies have targeted dorsally accessible brain structures like the inferior colliculus (IC) and auditory cortex (AC) in awake transgenic mice that had been exposed to anesthetics during preparatory surgery.
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FIGURE 4 | Time-dependent inhibition of Isoflurane. (A) Schematic diagram of the pharmacological protocol for testing Isoflurane inhibition. The first step was to record the spontaneous activity during 10 min under K/X mix (black box). Then we applied Isoflurane 1.5% for 90 s followed by two recording rounds of 10 min to obtain MUA after 10 or 20 min, respectively (red boxes). After that, ISO was turned off to evaluate the washout for the next 20 min. (B) Three examples showing the ISO inhibition and its washout at different developmental ages (P3, P6, and P9). Normalized MUA frequency relative to the K/X mix state (Norm MUA freq). (C) Percentage of Inhibition (MUA frequency under ISO/MUA frequency under K/X mix) after 10 (left) or 20 min (right) with ISO at different developmental stages. Statistical analysis show that ISO at 10 min had a greater inhibition at P6 in comparison with P12 (K-W, p = 0.007; multiple comparisons, * indicates multiple comparisons, p < 0.05; n = 3).

(Babola et al., 2018, 2020). However, if ventrally accessible brainstem structures are targeted to perform imaging and/or electrophysiological recordings, a general anesthetic must be used to perform recordings in animals that are mechanically ventilated (Rodríguez-Contreras et al., 2014; Di Guilmi et al., 2019), or that are immobilized to facilitate accurate electrode targeting (Sonntag et al., 2009; Clause et al., 2014). In this work we studied the impact of K/X mix and ISO on the spontaneous electrical activity of medial superior olivary neurons in neonate rodents.

This study provides three main findings that are relevant to understand the role of medial superior olivary neurons in auditory system development. First, to the best of our knowledge, this is the first study that compares K/X mix and ISO in the same rodent species. The firing pattern of single units recorded in K/X mix was similar to that reported in ISO anesthetized rat pups (Figure 1; Tritsch et al., 2010). Furthermore, this is the first study that identified burst-firing neurons outside the MNTB, which due to their location were mapped to the medial portion of the SOC (Figure 2). Second, it was found that ISO displayed a large scatter on its suppressing effects on electrical activity of auditory brainstem neurons. In pups that were pre-anesthetized with K/X mix, around two thirds of recorded cells were inhibited by 1.5% isoﬂurane, the isoﬂurane EC50 estimated under controlled conditions (Figures 1, 4; Wang Y. et al., 2020). However, an estimated one third of recorded units, all of which had a regular firing pattern, were resistant to 1.5% and were inhibited by 5% isoﬂurane delivered over 20 min. Third, this work showed that electrical ensemble activity of medial superior olivary neurons ramps up during development before hearing onset, with a similar developmental proﬁle under both anesthetics (Figure 3).

One caveat of the present study is that we were not able to study the effect of different concentrations of K/X mix on auditory brainstem neuronal activity. Future studies could first induce and maintain an anesthetic state with ISO, while K/X mix is delivered and titrated with precision via intravascular infusion. Another limitation of the present study is that the spatial location of neurons with different firing properties was inferred after intense post hoc histological analysis, and the functional properties of those cells could not be studied in the same animal. We consider that our results shed light on the use of anesthetics as tools for future studies that enable patch-clamp electrophysiology combined with imaging studies of medial superior olivary neurons in vivo. The approach would be to use K/X mix to induce a stable anesthetic state and ISO to probe
neuron inhibition on burst and regular firing neurons, which are predicted to have different responses to ISO. A combined optical calcium reporter imaging and electrophysiology approach would be desirable and useful to study MOC neurons, whose cell bodies are diffusely located in the ventral nucleus of the trapezoid body (VNTB) (Warr, 1975; Warr and Guinan, 1979; Guinan et al., 1983) and could be labeled with fluorescent reporters using genetic approaches (Cadenas et al., 2020).

Lastly, it is important to consider how the present study may advance our understanding of the mechanisms of anesthetic action throughout life, since neurons in the medial superior olive have also been studied in adult rodents anesthetized with K/X mix to investigate auditory processing (Lorteije et al., 2009; van der Heijden et al., 2013). It is noted that the K/X mix and ISO have different synaptic targets [i.e., postsynaptic NMDA receptors (MacDonald et al., 1987) or presynaptic effect (Wang Y. et al., 2020)]. In agreement with previous reports, synaptic depressant percentage depended on ISO concentration and the developmental age of pups (Wu et al., 2004; Wang Y. et al., 2020). Different reasons related to the plastic changes of auditory synapses (Wu et al., 2004), the timing of hearing onset (Qi et al., 2020), and the fact that the principal target of ketamine is the NMDA receptor (MacDonald et al., 1987) which declined to very low levels at the end of the second postnatal week (Taschenberger and von Gersdorff, 2000; Joshi and Wang, 2002), may explain the results of this study. Inhaled anesthetics diffuse from the alveoli into arterial and capillary blood, and are assumed to equilibrate rapidly with the well-perfused central nervous system (Hemmings et al., 2005). Thus, the heterogeneity of inhibitory effects observed in the results of this study, may also be explained by the hypothesis that the physical and functional properties of the vasculature, the diffusive properties of the extracellular space between vessels and neurons, and the clearance mechanisms of extracellular fluid available during this developmental window may be important factors that determine the access of anesthetics to target neurons in the brain (Nicholson et al., 2000; Shi and Rodriguez-Contreras, 2016; Mestre et al., 2020). A major difference in the effects of ISO in adult animals compared to neonates is the induction of slow waves of neuronal activity, an effect not described in neonates (Golshani et al., 2009; Brier et al., 2019). In this study, we observed that a group of recorded cells were inactive. One of the responsible factors of neuronal excitability is the resting membrane potential (Vm). Volatile anesthetics induced negative potentials on cortical cells (Petersen et al., 2003) and the hyperpolarization of postsynaptic MNBT neurons in brain slices (Wang Y. et al., 2020). Rusu and Borst (2010) demonstrated in slices containing the MNBT that Vm decreases during the first postnatal week becoming progressively more negative. During the first two postnatal weeks, the heterogeneity in the intrinsic properties maturation is larger than in adulthood, and probably it is a substrate for our observations. However, the effect of ISO on the Vm of these cells is under debate (Wu et al., 2004; Wang Y. et al., 2020) and a further study of molecular mechanisms during development is necessary to clarify the results.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

ETHICS STATEMENT

The animal study was reviewed and approved by the Institutional Animal Care and Use Committee of the City College of New York.

AUTHOR CONTRIBUTIONS

AR-C designed and performed the patch-clamp, multielectrode, and histology experiments. MD designed and performed the multi-electrode experiments. MD and AR-C analyzed the data and wrote and edited the manuscript. Both authors contributed to the article and approved the submitted version.

FUNDING

This work was supported by the National Institutes of Health Grant SC1DC015907 to AR-C. MD received the Company of Biologists traveling fellowship DEVTF-160703 and a “Bec.Ar” fellowship for a short stay supported by the Argentinian Government.

ACKNOWLEDGMENTS

We would like to acknowledge the Science Division Core Imaging Facility and Dr. Jonathan Levitt for providing access to a microtome to conduct histology procedures.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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