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Brainstem control of vocalization and its coordination with respiration

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1	Brainstem control of vocalization and its coordination with respiration
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Phonation critically depends on precise controls of laryngeal muscles in coordination with ongoing 34 respiration. However, the neural mechanisms governing these processes remain unclear. We 35 identified excitatory vocalization-specific laryngeal premotor neurons located in the 36 retroambiguus nucleus (RAm^{VOC}) in adult mice as both necessary and sufficient for driving vocal-37 cord closure and eliciting mouse ultrasonic vocalizations (USVs). The duration of RAm^{VOC}-38 activation can determine the lengths of both USV syllables and concurrent expiration periods, with 39 the impact of RAm^{VOC}-activation depending on respiration phases. RAm^{VOC}-neurons receive 40 inhibition from the preBötzinger complex, and inspiration-needs override RAm^{VOC}-mediated-41 vocal-cord closure. Ablating inhibitory synapses in RAm^{VOC}-neurons compromised this 42 inspiration gating of laryngeal adduction, resulting in discoordination of vocalization with 43 respiration. Our study revealed the circuits for vocal production and vocal-respiratory coordination. 44

45 **One-Sentence Summary:**

Identification of RAm^{VOC}-neurons as the critical node for vocal production and vocal-respiratory
 coordination.

Vocalization plays essential roles in communication in many species (1, 2). While the 48 complexity of vocalization (i.e. articulation) varies depending on species, the fundamental sound 49 production process (i.e. phonation) shares similarities. Phonation process dominantly occurs 50 during expiration: narrowing of the larynx (vocal cord adduction) while simultaneously exhaling 51 air (3). In general, phonations do not happen during inhalation because inspiration requires opening 52 of the larynx (vocal cord abduction) (4). Furthermore, the need for inspiration suppresses 53 vocalization (breathing primacy), as everyday experience illustrates that we have to stop talking 54 when we need to breathe. Inappropriate adduction or abduction of the larynx in the wrong 55 respiration phases can lead to inspiration problems or hoarse vocalizations (5, 6). However, the 56 neural circuits that seamlessly coordinate laryngeal movements with respiration to produce 57 phonations and to prioritize breathing needs have yet to be clearly delineated. 58

59 We reasoned that the key to answer this question is to first identify the neurons that drive laryngeal adduction for vocalization, followed by determining their interaction with respiratory 60 circuits. The hindbrain contains premotor neurons that can activate laryngeal adductor 61 motoneurons (1, 2, 7). The nucleus retroambiguus (RAm) located in the caudal-ventral brainstem 62 63 is one key node for vocal production. Vocalizations induced by electrical stimulation of the midbrain periaqueductal gray (PAG) in decerebrate cats (8, 9) and anesthetized rats (10) are 64 65 suppressed by lesions of the RAm. Pharmacological and electrical stimulation of the RAm evokes elementary sounds (9-11), although such sounds do not resemble species-typical vocalizations. 66 The RAm region has vocalization-related neural activity (12), and shows a positive correlation 67 between unit activity and vocal loudness (13). Neural tracers injected in the RAm labels axonal 68 69 projections to the nucleus ambiguus (NA) where laryngeal motoneurons are located (14). However, the RAm region does not have anatomical demarcations and contains heterogeneous types of 70 neurons including neurons modulating respirations and other orofacial movements (15). Thus, it 71 remains unknown which populations in the RAm are vocalization-specific laryngeal premotor 72 73 neurons and whether they are necessary and sufficient to drive vocal cord adduction and phonation, and if so, how these neurons interact with respiratory circuit to ensure vocal-respiration 74 75 coordination and breathing primacy. With regard to respiration, intensive studies have been conducted on the inspiration rhythm generator, the preBötzinger complex (preBötC) (16-19). 76 However, only one study investigated the function of the preBötC during vocalizations in awake 77 animals (20). As such, it is still unclear how inspiration gates the activity of hindbrain vocal 78 production circuits. 79

We used mouse ultrasonic vocalization (USV) as a model system. During interactions with 80 female mice, male mice readily emit USVs comprising a string of syllables periodically interrupted 81 by inspiration, also called courtship songs (21, 22). Unlike audible vocalizations, which are 82 produced by air vibrating the tightly-closed vocal cords (23), USVs are produced by a whistle-like 83 mechanism: a jet stream of air coming through a small hole formed between the adducted vocal 84 cords (24-26), thereby generating pure-tone sounds in ultrasonic frequency range. Despite the 85 unique phonation mechanism, USVs still require laryngeal adduction and necessitate the adduction 86 occurring during expiration (24), thereby providing us a suitable model for vocal-respiratory 87 coordination. 88

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90 Vocalization-specific laryngeal premotor neurons in the brainstem

91 The activity of laryngeal muscles and motoneurons is controlled by premotor neurons in 92 the hindbrain (1, 7). However, the location and identity of the vocal premotor circuits in *adult*

mammals have yet to be revealed. We applied three-step monosynaptic rabies virus tracing (27) 93 (Fig. 1A), combining AAVretro-Cre (injected into laryngeal muscles in juvenile animals), Cre-94 dependent helper AAVs (to express TVA receptor and optimized rabies glycoprotein (oG) in 95 motoneurons), and pseudo-typed G-deleted rabies virus (EnvA^{M21}-RV-GFP, injected into the NA 96 in adults). Cre⁺ motoneurons were found around the NA (Fig. 1B), and trans-synaptically labeled 97 laryngeal premotor neurons were mostly observed in the brainstem (Fig. 1C), specifically in the 98 Kölliker-Fuse (KF), parvocellular reticular formation (PCRt), lateral paragigantocellular nucleus 99 (LPGi), intermediate reticular nucleus (IRt), preBötC, nucleus tractus solitarii (NTS), and RAm. 100 We registered all labeled neurons in the Allen common coordinate frame for the mouse brain 101 (Allen CCF) (28) and compared the map of laryngeal premotor neurons to our previously identified 102 103 maps of jaw and tongue premotor neurons (27) (fig. S1). The overall spatial distributions of laryngeal premotor neurons from different mice (n=3) were similar, but they were distinct from 104 those of jaw and tongue premotor maps (fig. S1). Labeled premotor neurons also had extensive 105 collateral projections to other branchial motor nuclei, including the trigeminal (5N), the facial (7N), 106 and the hypoglossal (12N) nuclei (fig. S1), suggesting that laryngeal premotor neurons might 107 simultaneously recruit other orofacial motoneurons for vocalization and perhaps for other orofacial 108 movements. 109

Previous studies have suggested that the RAm is a critical node for vocal production (7, 110 14). When we examined Fos mRNA expression (a marker for activated neurons) in male mice 90 111 min after female-induced courtship USVs (Fig. 1D), we detected robust Fos signals in the RAm 112 (fig. S2). By contrast, fewer and weaker *Fos* expressions were found in other hindbrain areas, 113 114 such as the preBötC in the same samples (fig. S2). Our laryngeal premotor tracing consistently labeled a cluster of RAm neurons (Fig. 1C). We further confirmed that the majority of rabies-115 traced laryngeal premotor neurons in the RAm induced Fos expression after bouts of courtship 116 USVs (68.6±13.1 %, GFP+ and Fos+ neurons/GFP+ neurons, n=4 mice, Fig. 1E). 117

We used the Fos-based cell targeting method called CANE (29) to label courtship USV-118 activated RAm neurons in male mice (RAm^{VOC}-neurons) (Fig. 2A). After expressing GFP in 119 RAm^{VOC}-neurons via CANE, we re-exposed male mice to females to re-elicit USVs and Fos 120 expression and confirmed that labeled RAm^{VOC} were indeed Fos+ (Fig. 2B). We further registered 121 the locations of all CANE-captured RAm^{VOC}-neurons in the Allen CCF and confirmed that their 122 positions overlapped with those of the rabies-traced RAm laryngeal premotor neurons (Fig. 2C). 123 We further examined the expression of ChAT, a molecular marker for motoneurons, and found 124 that none of the labeled RAm^{VOC}-neurons expressed ChAT (Fig. 2D), i.e., CANE did not capture 125 cholinergic motoneurons. Furthermore, the axonal boutons from RAm^{VOC}-GFP cells innervated 126 ChAT positive motoneurons around the NA (Fig. 2D), consistent with them being vocal premotor 127 neurons. Lastly, in-situ hybridization using Vglut2 and Vgat probes showed that majority of 128 RAm^{VOC}-neurons were glutamatergic (Vglut2+/RAm^{VOC}: 85.1±0.1%, Vgat+/RAm^{VOC}: 12.9±0.1%, 129 n=3 mice, Fig. 2E), suggesting that they provide excitatory inputs to laryngeal motoneurons. 130

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Silencing RAm^{VOC}-neurons abolishes both ultrasonic and audible vocalizations

To dissect the functional role of RAm^{VOC}-neurons, we bilaterally expressed tetanus toxin light chain (TeLC) to inhibit their synaptic outputs (*30*) or expressed GFP as controls using CANE (Fig. 2F). RAm^{VOC}-GFP male mice emitted robust USVs in the presence of female mice before and after CANE-mediated expression (Fig. 2G upper and H left). In contrast, RAm^{VOC}-TeLC mice failed to vocalize in response to female mice after TeLC expression (Fig. 2G bottom and H right). The effect of silencing RAm^{VOC}-neurons was robust and consistent: all six RAm^{VOC}-TeLC mice
 had complete mutism during courtship (Fig. 2I).

In addition to social USVs, mice also elicit audible squeaks in response to strongly aversive 140 stimuli (31). Prior studies suggested that USVs and squeaks are triggered by different neural 141 pathways (31, 32). For example, a recent study showed that inhibition of the PAG-RAm pathway 142 only abolished USVs but not pain-elicited audible vocalizations (32). We evoked squeaks in mice 143 using a tail-pinch stimulus (Fig. 2J). While control RAm^{VOC}-GFP mice responded with robust cries, 144 RAm^{VOC}-TeLC mice were silent (Fig. 2K and L). Furthermore, when we applied foot-shocks, 145 RAm^{VOC}-GFP (Movie S1), but not RAm^{VOC}-TeLC mice (Movie S2), squeaked, even though all 146 mice exhibited escape behaviors, indicating that nociceptive responses of the RAm^{VOC}-TeLC mice 147 were intact. 148

To rule out the possibility that mutism in the RAm^{VOC}-TeLC mice originated from general breathing abnormalities, we habituated mice on a treadmill wheel and gently encouraged them to run (fig. S3). Running changes both the frequency and amplitude of breathing in mice (*33*). The modulation of respiration by running in RAm^{VOC}-TeLC mice remained intact as that in the control group (RAm^{VOC}-TeLC (n=3) vs RAm^{VOC}-GFP (n=4). Changes in inspiratory amplitude: 27.8±8.4% vs 24.4±2.6%, p=0.8597; expiratory amplitude: 12.6±6.6% vs 6.5±1.1%, p=0.5959; frequency: 36.7±18.3% vs 27.3±9.5%, p=0.5959, Mann-Whitney U test, fig. S3).

We also observed some axon collaterals of RAm^{VOC}-neurons in the thoracic spinal cord 156 segment (fig. S4), where abdominal spinal motor neurons for active expiration are located, 157 suggesting that RAm^{VOC} might be involved in increasing expiratory activity needed for generating 158 sound (phonation). To test this idea, we measured abdominal EMG of anesthetized RAm^{VOC}-TeLC 159 mice during PAG stimulation-induced vocalizations (fig. S4). A previous study has shown that 160 optogenetic stimulation of RAm-projecting PAG neurons (PAG^{RAm}) could reliably elicit USVs in 161 mice (32). PAG^{RAm} neurons were labeled by injecting AAVretro-FlpO in the RAm, and injecting 162 Flp-dependent optogenetic activator ChRmine (34) in the PAG, and in the same male mouse, 163 RAm^{VOC} neurons were targeted to express either GFP or TeLC using CANE (fig. S4). While 164 PAG^{RAm} stimulation reliably elicited abdominal EMG activity concurrent with USVs in the GFP 165 control mice, the same stimulation failed to elicit USVs and abdominal EMG responses in the 166 TeLC mice (fig. S4). 167

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169 *RAm^{VOC}-activation is sufficient to elicit and modulate USVs in mice*

In addition to active expiration, vocal production critically depends on vocal cord 170 adduction. The nearly closed larvnx is essential for the exhaling jet stream of air to whistle USVs, 171 or to vibrate the vocal cords to produce audible sounds (24-26). To determine whether RAm^{VOC}-172 neurons are sufficient to close the vocal cords and elicit USVs, we expressed ChRmine (34) in 173 these neurons using CANE in male mice (Fig. 3A). First, the larynx was imaged with a camera 174 while mice were anesthetized and placed in a prone position (Fig. 3B). The vocal cords naturally 175 widened and narrowed (but not fully closed) rhythmically (Fig. 3C, Movie S3), in phase with 176 inhalation and exhalation, resulting in periodic changes in the size of the glottal area (Fig. 3D). 177 Optogenetic activation of RAm^{VOC} with 5s continuous laser illumination instantaneously closed 178 the vocal cords, and the laryngeal adduction persisted throughout the stimulation (Fig. 3D, n=3 179 mice, Movie S3). This prolonged laryngeal adduction was interrupted by occasional glottal 180 openings during the 5s stimulation in all mice tested (this point is further elaborated below). We 181 next stimulated RAm^{VOC} in awake male mice to check whether this was sufficient to elicit USVs 182

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(Fig. 3E). Applying a brief 100ms laser pulse reliably induced USVs time-locked to each pulse 183 (Fig. 3F). The onset latencies of the optogenetic-induced ultrasonic vocalizations were short 184 (39.0.0±1.1ms, Fig. 3G). All RAm^{VOC}-activation-elicited vocalizations were in ultrasonic range 185 (RAm^{VOC}-USV), and the syllable patterns of RAm^{VOC}-USVs included several typical types of 186 female-directed USVs (35) (up, step-down, chevron, two-steps, short, but also unstructured ones, 187 Fig. 3H). We also compared RAm^{VOC}-USVs and female-directed USVs for several acoustic 188 features, and observed similar distributions for loudness, spectral purity, and pitch variance (Fig. 189 3I). Note that the mean frequency of the RAm^{VOC}-USVs was different, i.e., lower than that of the 190 female-directed USVs in the same mice (RAm^{VOC}-USVs: 61.8±0.4 kHz, female-directed: 191 79.6 \pm 0.2 kHz, p \leq 0.0001, Mann-Whitney U test), indicating other neurons are needed for 192 193 producing the full frequency range of natural USVs.

Given that a brief RAm^{VOC} -activation elicited a single short USV syllable (Fig. 3F), we also tested whether RAm^{VOC} -activation can alter the length of individual USV syllables. We varied the duration of optogenetic stimulation of RAm^{VOC} (50, 100, and 200ms), and observed that indeed the length of RAm^{VOC} -USV syllables were proportionally correlated to the duration of laser stimuli (Fig. 4B and D).

199

200 *Vocalization-respiration coordination during RAm^{VOC} activation*

For normal vocalization, sound is exclusively produced during the expiration phase (4). 201 The results described above highlighted the role of RAm^{VOC}-neurons in driving laryngeal 202 adduction while coordinating expiration efforts. However, inspiration-needs must be prioritized 203 (breathing primacy) to ensure survival. To investigate the precise role of RAm^{VOC} in vocal-204 respiration coordination, we simultaneously measured USVs and respiratory activity in awake 205 mice while optogenetically stimulating RAm^{VOC} with different durations (50, 100, and 200ms) 206 (Fig. 4A-C). Longer RAm^{VOC}-activation induced longer duration of expiration characterized by a 207 flat period on the respiratory traces (Fig. 4C and E). The durations of RAm^{VOC}-induced-USVs and 208 flat expirations were highly correlated (R²=0.922), consistent with the notion that RAm^{VOC}-209 activity coordinately mediates vocal cord closure and expiration. 210

We next asked whether the impact of RAm^{VOC}-activation is dependent on the current on-211 going respiratory phases. To test this idea, we analyzed the latencies and durations of RAm^{VOC}-212 induced flat expirations and USV syllables with respect to the onsets of laser RAm^{VOC}-activation 213 in respiration phases (Flaser, Fig. 4F). Interestingly, RAm^{VOC}-stimulation at the early expiration 214 (F_{laser} during 0 to 0.5 π) and late inspiration phases (F_{laser} during -0.5 π to 0) produced longer 215 durations of expirations and USVs with short latencies, while RAm^{VOC}-activation in the late 216 expiration (F_{laser} during 0.5π to π) and early inspiration phases (F_{laser} during $-\pi$ to -0.5π) elicited 217 shorter expirations and shorter USVs with longer latencies (Fig. 4 G and H). 218

With 200ms of RAm^{VOC}-activation, we occasionally observed a full inspiration cycle 219 during stimulation (200ms, Fig. 4C). Similarly, in the anesthetized larynx imaging preparation, the 220 vocal cords were occasionally open during prolonged 5s RAm^{VOC}-activation, presumably due to 221 an "override" by the need for inspiration (Fig. 3D). To further investigate this inspiratory gating 222 of vocalization/vocal adduction in awake mice, we applied 2s continuous RAm^{VOC}-activation. This 223 2s stimulation produced multiple USV syllables accompanied by concurrent flat expiration 224 periods, which were periodically interrupted by intervening inspirations (Fig. 4I). The amplitudes 225 of the intervening inspirations were similar to those in the baseline conditions, indicating that these 226 are normal breaths (Fig. 4I). We projected the onsets and offsets of the multiple USV syllables 227

evoked by the 2s RAm^{VOC}-activation onto respiration phase maps (Inspiration: $-\pi$ to 0, Expiration: 0 to π , Fig. 4J). All syllables were exclusively found in the expiration phase (Fig. 4K), consistent with the notion that intervening inspirations can stop the on-going USVs evoked by RAm^{VOC}activation, i.e., inspiration gates and sets the basic rhythm of vocalization.

232

233 Inhibitory inputs to RAm^{VOC} are essential for inspiration gating of vocalizations

We hypothesized that inhibitory inputs onto RAm^{VOC}-neurons are the key for the periodic 234 suppression of vocalization by inspiration. To identify the source of inspiration-related inhibitory 235 inputs to the RAm^{VOC}-neurons, we performed monosynaptic tracing of presynaptic neurons to 236 RAm^{VOC} (preRAm^{VOC}). This was achieved by expressing TVA and oG in RAm^{VOC} using CANE, 237 followed by infecting these neurons with EnvA^{M21}-RV-GFP (Fig. 5A). Tracing results showed that 238 RAm^{VOC}-neurons receive excitatory inputs from the PAG, the parabrachial (PB)/KF, and other 239 areas (Fig. 5B). Excitatory PAG neurons are known to be required for eliciting USVs but not for 240 generating rhythmic vocal patterns (32). The dominant source of inhibitory inputs to RAm^{VOC}-241 neurons was the preBötC (Fig. 5B), the inspiration rhythm generator (19). In our mapping of 242 laryngeal premotor neurons, we also labeled a population of inhibitory neurons in the preBötC 243 (fig. S5). Thus, the preBötC provides inhibitory inputs to both vocal motoneurons (MN^{VOC}) and 244 to RAm^{VOC} (Fig. 5C), consistent with a recent axonal tracing study of inhibitory preBötC neurons 245 (36). These results suggest that the inspiration-controlled periodic patterns of USVs could be 246 generated by tonic excitatory inputs from the PAG to RAm^{VOC} to induce vocal cord adduction (and 247 concurrent expiration), which is gated by rhythmic inhibition from the preBötC to both MN^{VOC} 248 and RAm^{VOC} (Fig. 5C). 249

To validate the functional relevance of the anatomical connections identified above, we 250 decided to block inhibitory inputs to RAm^{VOC}-neurons. Based on the circuit diagram, we predicted 251 that disinhibited RAm^{VOC} would provide stronger and tonic excitatory drive to MN^{VOC}, that 252 counters the rhythmic inhibitory drive from the preBötC, such that vocal cord adduction may 253 happen even during inspiration. Furthermore, if the activity of disinhibited RAm^{VOC} was 254 sufficiently elevated, spontaneous vocalization (in the absence of social interactions) might occur. 255 We expressed GFE3 in glutamatergic RAm^{VOC}-neurons using CANE (RAm^{VOC}-GFE3 mice), with 256 RAm^{VOC}-GFP mice as control (Fig. 5D). This was achieved by injecting Cre-dependent CANE-257 hSyn-DIO-tTA together with AAV-TRE3G-GFE3 (or GFP) in the RAm in Fos^{TVA}/Vglut2-Cre 258 double transgenic male mice after bouts of courtship USVs. GFE3 is a ubiquitin ligase specifically 259 targeting the inhibitory post-synaptic scaffolding protein gephyrin for degradation (37), thereby 260 reducing phasic synaptic inhibition onto RAm^{VOC}-neurons. To reliably elicit USVs in awake head-261 fixed mice, we again chose to perform optogenetic stimulation of RAm-projecting PAG neurons 262 (PAG^{RAm}) (32). Briefly, in the same RAm^{VOC}-GFE3 or control mice, we also expressed ChRmine in RAm-projecting *Vglut2*⁺ PAG neurons (PAG^{RAm/vglut2}) using a Flp/Cre intersectional strategy 263 264 (Fig. 5D). In control RAm^{VOC}-GFP mice, continuous pulses of optogenetic stimulation of 265 PAG^{RAm/vglut2} reliably elicited USVs but only during expirations, as the expirations were 266 periodically interrupted by the inspiration flows (Fig. 5E-F upper panels). In addition, the peak 267 flow values for the inspiration (downward trace) increased during the optogenetic PAG stimulation 268 (123.1±6.1%, n=4 mice, Fig. 5E and G), suggesting PAG^{RAm/vglut2} activation enhances inspiration 269 (likely for inhaling sufficient air for vocalization). By contrast, in RAm^{VOC}-GFE3 mice, the 270 inspiratory interruption of vocalization was severely compromised during continuous 271 PAG^{RAm/vglut2} activation (Fig. 5E, lower panels). The amplitude of the few intervening inspirations 272 during PAG stimulations was significantly reduced compared to the average inspiration peak 273

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before stimulation (49.6±10.5%, n=5 mice, p=0.020, Mann-Whitney U test for GFE3 vs GFP mice, 274 Fig. 5F and G, lower panels). We observed that asthma-like vocal sounds were produced during 275 the inspiration periods in RAm^{VOC}-GFE3 (21.8±5.4%, n=5 mice, Fig. 5F, gray-shaded region, and 276 5G), while these abnormal inspiratory vocal sounds were never observed in the RAm^{VOC}-GFP 277 control mice during PAG^{RAm/vglut2} activation. Thus, removing inhibitory synaptic inputs to 278 RAm^{VOC}-neurons compromises inspiration-gating of vocalization. The reduced inspiration 279 amplitude is likely caused by persistent vocal cord adduction, due to a tonic excitatory drive from 280 the disinhibited RAm^{VOC}. This persistent vocal cord adduction during inspiration could also 281 explain the abnormal asthma-like inspiratory vocalizations. Finally, consistent with the idea that 282 tonic activation of disinhibited RAm^{VOC}-neurons would cause spontaneous vocal cord closures, 283 RAm^{VOC}-GFE3 mice also produced occasional spontaneous USVs in the absence of social 284 contexts $(0.5\pm0.2 \text{ VOC/s}, n=6 \text{ mice}, \text{ fig. S6})$, whereas control male mice almost never utter 285 spontaneous USVs. 286

287

288 Discussion

We detected a vocalization-specific laryngeal premotor population in the RAm region of 289 the caudal hindbrain (RAm^{VOC}) as the critical node for driving laryngeal adduction and phonation. 290 We further uncovered neural mechanisms involving preBötC-RAm^{VOC} interactions that ensure 291 breathing primacy by allowing rhythmic inspirations to pace vocalizations. It has been debated 292 whether the neural circuits for laryngeal adduction and vocal production are distributed across the 293 ventral brainstem (7) or localized in one small area, such as the RAm (14). Here we found that 294 inhibition of RAm^{VOC}-neurons not only abolished USVs in social contexts but also audible squeaks 295 during aversive states (tail-pinch or foot-shock). Thus, RAm^{VOC} represents a singular necessary 296 locus for all phonations. On the other hand, optogenetic stimulation of RAm^{VOC}-neurons was 297 sufficient to produce and only produced USVs, but not audible sounds. USVs and squeaks in 298 rodents have different acoustic features. USVs lie above ultrasonic range (> 20kHz) with pure 299 tones (21, 22), and rodents use aerodynamic mechanisms to produce USVs (24-26), while audible-300 squeaks occupy a human hearing frequency range (below 20kHz), with harmonics (38). Thus, 301 squeaks likely require additional circuit elements, such as those driving strong air exhalation, 302 which are not activated or recruited by RAm^{VOC}. 303

USVs can be further modulated in terms of frequency and duration. The duration of mouse 304 vocalizations could be modulated by RAm^{VOC}-activity, but the mean frequency of RAm^{VOC}-USVs 305 were lower than those of female-directed USVs in the same animals (Fig. 3). These data suggest 306 that another parallel premotor pathway to larvngeal motor neurons (e.g., to vocal tensor muscles, 307 such as cricothyroid muscles) might be involved in vocal frequency regulation. One potential 308 frequency modulating region is the PCRt, which contains larvngeal premotor neurons as shown in 309 310 our transsynaptic tracing study (Fig. 1C). This region, referred to as the vocalization-related parvicellular reticular formation (VoPaRt) in rats, is a node for high frequency vocalization (10). 311 For duration modulation, we showed that optogenetically increasing the time of RAm^{VOC}-312 activation elongated the syllable length (Fig. 4B and D). Interestingly, transsynaptic tracing of pre-313 RAm^{VOC} neurons labeled inputs in the PB/KF (Fig. 5B), which could be the endogenous region 314 controlling RAm^{VOC}-activation and vocal duration based on previous pharmacological studies 315 (39). However, the PB/KF regions are heterogeneous, including intermingled non-vocal 316 respiratory neurons (40, 41), therefore future work targeting vocal-specific PB/KF will be needed 317 to reveal the precise role of PB/KF in controlling vocal durations. Furthermore, it will be 318

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interesting to know whether and how the other recently identified brainstem vocal modulatory loci, the iRO in neonate mice (42) interacts with RAm^{VOC} to modulate other features of vocalizations.

Breathing is vital for survival. As breathing and vocalization both occur in the airway, 321 laryngeal closure for sound production needs to be precisely controlled and coordinated with 322 respiration. Failure in such coordination could lead to vocal cord dysfunction and breathing 323 problems (5, 6). We found evidence of inspiration dominance over RAm^{VOC}-USVs: the effect of 324 brief RAm^{VOC}-activation was delayed and attenuated around the onset of inspirations: USV 325 syllables produced by prolonged-RAm^{VOC}-activation were periodically interrupted by full 326 inspiration peaks (Fig. 4). We found that the inspiration rhythm generator preBötC, where Vgat+ 327 and GlyT2 + neurons are found (36, 43), provides the main source of inhibitory inputs to RAm^{VOC} 328 (Fig. 5). Chronic disinhibition of RAm^{VOC} in RAm^{VOC}-GFE3 experiments reduced the amplitudes 329 of inspiratory gating during vocalization, produced hoarse sound in inspiration phases as well as 330 331 spontaneous USVs in the absence of social context (Fig. 5 and fig. S6). Taken together, our results support a conceptual model (Fig. 5C) in which the timing of phonation is controlled by the 332 combined activity of preBötC and RAm^{VOC}, with inspiration playing a dominant role in setting the 333 basic rhythm of vocalization, while RAm^{VOC} driving vocal cord closure and modulating syllable 334 durations within the limit set by inspiration. This mechanism produces the periodic alternating 335 patterns of vocalization and inspiration. In human speech, multiple syllables can be uttered within 336 one breath, and in that case, a separate multi-syllable rhythm generator within expiration period 337 might be needed. We also labeled laryngeal premotor neurons in NTS (Fig. 1C), which is a region 338 receiving inputs from vagal pulmonary afferents (43). It is possible that the pulmonary-NTS 339 pathway is involved in the transition between inspiration and vocalization (44). When the lungs 340 are inflated with enough air, this pathway may help to inhibit the activity of preBötC and facilitate 341 the transition to vocalization and expiration. Future work should test whether the pulmonary-NTS 342 circuit represents the third node in modulating vocal patterns. 343

Finally, we want to point out that our study focused only on the "phonation", but not the 344 complex "articulation" aspect of vocalization. Vocal articulations are among the most complicated 345 motor patterns generated by humans (and many mammals) as they require coordinated control of 346 the laryngeal, facial, tongue, jaw, and respiratory muscles. How this is achieved remains poorly 347 understood. In our transsynaptic tracing studies, we labeled a large population of neurons in the 348 reticular formation, and we found that larvngeal premotor neurons also project to other orofacial 349 motor nuclei (fig. S1). However, the identities of these premotor neurons are unknown, and more 350 work will be needed to determine whether and how these neurons are involved in complex 351 articulations. 352

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480 **Author contributions:**

- F.W. and J.P conceptualized the project, designed experiments, and wrote the paper with input
 from all authors. J.P. performed the majority of experiments. J.P. and J.T. analyzed data. S.C.
 performed histology works. S.Z., A.H., and J.T. produced key viral vectors used in this study.
 B.H. and S.C. provided animal husbandry. F.W. supervised all the work.
- 485 **Competing interests:**
 - The authors declare no competing interests.

487 **Data and materials availability:**

- All data and code are available in the main text, the supplementary materials, or Dryad (https://doi.org/10.5061/dryad.vmcvdnd0m).
- 490 Supplementary Materials
- 491 Materials and Methods
- 492 Figs. S1 to S6
- 493 Movies S1 to S3
- 494 References (46-48)

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Fig. 1. Transsynaptic mapping of laryngeal premotor neurons and vocalization-induced Fos activity in the RAm.

(A) A schematic for three-step monosynaptic rabies virus strategy using AAVretro-Cre, helper 511 virus (AAV-Flex-oG, AAV-Flex-TVA-mCherry), and monosynaptic rabies virus (EnvA^{M21} 512 coated) to map laryngeal premotor neurons. (B) Laryngeal motoneurons (red) labeled by 513 AAVretro-Cre in the brainstem of an Ai-14 reporter mouse. (C) Laryngeal premotor neurons 514 (green) in the KF, PCRt, LPGi, preBötC, IRt, VRG, NTS, and RAm. (D) A schematic for Fos (for 515 1h) or Fos mRNA (30min) induction experiments in a social-context eliciting USVs in male mice. 516 (E) Laryngeal premotor neurons (green) and Fos (magenta) labeling in the RAm (upper). A 517 zoomed-image of the boxed area (bottom). Neurotrace Blue was used to visualize neuronal 518 structures. Scale bars, 200 µm (B, C, E upper); 50 µm (E bottom). 519



524 Fig. 2. Vocalization-induced Fos positive neurons in the RAm (RAm^{VOC}-neurons) are 525 excitatory laryngeal premotor neurons and required for vocalization in mice.

(A) A schematic for CANE experiments to capture vocalization-induced Fos positive neurons in 526 the RAm. (B) RAm^{VOC}-neurons (green) with Fos immunolabeling (red). (C) RAm^{VOC}-neurons 527 (green) with the laryngeal premotor neurons (grey) in the Allen CCF in coronal (left) and sagittal 528 (right) views. (D) RAm^{VOC}-neurons (green) with ChAT immunolabeling (magenta). Left (soma) 529 and middle (axon terminals). The right panel highlights the NA region of the middle panel. (E) 530 RAm^{VOC}-neurons (green) with fluorescent in situ hybridization labeling for Vglut2 (magenta) 531 (left). Group data of Vglut2 and Vgat from n=3 mice. (F) A schematic for expressing TeLC in 532 RAm^{VOC}-neurons. (G) Spectrograms of female-directed USVs of RAm^{VOC}-GFP controls (upper) 533 and RAm^{VOC}-TeLC (bottom) mice. (H) USV rates of male mice during courtship behaviors for 10 534 min. Blue vertical lines indicate the time of female introduction (\mathcal{Q}). Grey and green plots for a 535 RAmVOC-GFP mouse and a RAm^{VOC}-TeLC mouse, before and 2 weeks after virus injection (left 536 and right, respectively). (I) The total numbers of USV syllables during 10min social interactions 537 (RAm^{VOC}-TeLC, green, n=6; and control, grey, n=3). (J) A schematic for recording tail pinch-538 induced audible squeaks. (K) Spectrogram (upper) and sound intensity plots (bottom) of audible 539 squeaks from RAm^{VOC}-GFP (grey, left) and RAm^{VOC}-TeLC (green, right) mice. Red vertical lines 540 indicate the onset of tail-pinch stimuli. (L) Average intensity of squeaks during tail-pinch (n=3 for 541 each group). 542

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556 Fig. 3 Optogenetic activation of RAm^{VOC}-neurons robustly elicits USV-like vocalizations in 557 mice.

(A) Schematic for expressing ChRmine to RAm^{VOC}-neurons using CANE method. (B) Schematic 558 for visualizing the vocal cords in anesthetized mice. (C) Images showing opened (left) and closed 559 (right) vocal cords. Red dots indicate the cartilage parts of the vocal cords that are used to track 560 the glottal area (red rectangle). (D) The response of the glottal area to RAm^{VOC}-opto-activation. 561 Green bar (5s) indicates the laser stimulation period. E) Schematic for recording vocalization of 562 awake mice in a head-fixed condition. F) Sound-time raw traces (upper) and corresponding 563 frequency-time spectrogram (bottom) during a train of brief laser pulses (laser wavelength = 564 560nm, 100ms of 4 pulses with 2s intervals). (G) Latency distribution of RAm^{VOC}-USVs (laser 565 duration: 100ms, 443 syllables, n=3 mice). (H) Examples of RAm^{VOC} (left upper row, red) and 566 female-directed USVs (left bottom row, grey). A single box spans 120ms (x axis) and 30 to 125kHz 567 (y axis). Classification results of RAm^{VOC}-USVs (right). (I) Distributions of four acoustic features 568 (loudness, spectral purity, mean frequency, and pitch variance) of RAm^{VOC}-USVs (443 syllables, 569 n=3 mice) and female-directed USVs (4960 syllables, n=3 mice). 570

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579 Fig. 4 RAm^{VOC}-activation can modulate the duration of USVs and concurrent expiratory 580 periods until interrupted by the need for breathing.

(A) Schematic for recording vocalization and respiration in RAm^{VOC}-ChRmine mice. (B) USV 581 syllables evoked by three different durations of RAm^{VOC} laser activation (50, 100, and 200ms). 582 (C) Respiratory responses to the RAm^{VOC}-activation (Left: 50ms, Right: 200ms). 13 trials are 583 aligned to the laser onsets and overlayed. Green lines indicate RAm^{VOC}-induced flat expiration 584 periods. (D) Average duration of RAm^{VOC}-USVs (n=3 mice). (E) Average duration of RAm^{VOC}-585 induced flat expiration periods (n=3 mice). (F) A schematic for defining laser stimulation phase 586 (Φ_{laser}) and latency and duration of RAm^{VOC}-induced flat expiration and USV to stimulation 587 (upper). Black trace indicates normalized airflow. Φ_{laser} is defined as a phase of laser onsets with 588 respect to the expected airflow (Inspiration: $-\pi$ to 0, Expiration: 0 to π). Four cases of different 589 stimulation onset phases (bottom). (G) Relationship between Φ_{laser} , and latency and duration of 590 RAm^{VOC}-expirations (rho, ms scale). The same-color solid lines represent polynomial-fitted lines. 591 Red and blue circle-arrows indicate the expiration (0 to π) and inspiration phases (- π to 0), 592 respectively. (H) Relationship between Φ_{laser} , and latency and duration of RAm^{VOC}-USVs. (I) 593 USVs (upper) and respiratory responses (bottom) to the 2s RAm^{VOC}-activation. Blue dots indicate 594 the inspiratory flow peaks. (J) Projection of onset and offset of a RAm^{VOC}-USV onto a respiratory 595 phase. (k) Phase density distribution of the onsets (red) and offsets (pink) of RAm^{VOC}-USVs. Blue 596 597 and red dash lines represent arbitrary inspiration and expiration phase, respectively.



Fig. 5 Ablating inhibitory synapses on RAm^{VOC}-neurons compromised vocal-respiratory coordination.

- 612 (A) Schematic for transsynaptically tracing preRAm^{VOC}-neurons (left). CANE and rabies labeled-613 source cells (magenta: TVA, cyan: GFP) in RAm (right). Dash-circles indicate RAm areas. (B)
- 613 source cells (magenta: TVA, cyan: GFP) in RAm (right). Dash-circles indicate RAm areas. (B) 614 preRAm^{VOC}-neurons (green) in the PAG, KF, and preBötC with in situ hybridization (magenta for
- 615 *Vglut2* and *Vgat*). (C) Schematic for the proposed neural mechanism for vocal-respiratory
- 616 coordination. (D) Schematic for ablating inhibitory synapses in RAm^{VOC}-neurons with GFE3
- 617 expression (RAm^{VOC}-GFE3), and concurrent expression of ChRmine in RAm projecting
- 618 glutamatergic PAG neurons. (E) Respiratory activities of the RAm^{VOC}-GFP (blue) and RAm^{VOC}-619 GFE3 (orange) mice in response to the PAG^{RAm/vglut2}-ChRmine stimulation for 2s. Blue dots
- 620 represent the inspiratory peaks. (F) Spectrogram (upper) with the respiratory responses (bottom).
- Grey bars label abnormal vocalizations in the inspiratory phases. (G) Average changes in the
- 622 inspiratory peaks of the mice (n=5, GFE3; n=4, GFP, upper) during the PAG^{RAm/vglut2} stimulation 623 over the baseline inspirations. The portions of the abnormal inspiratory vocalization among the
- 623 over the baseline inspirations. The portions of the abnormal inspiratory vocalization among the 624 $PAG^{RAm/vglut2}$ -induced vocalizations (n=5, GFE3; n=4, GFP, bottom). No inspiratory vocalization
- 625 was detected in the GFP control mice.

	Science
1 2	MAAAS
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4	Supplementary Materials for
5	
6	Brainstem control of vocalization and its coordination with respiration
/ 8 9	Jaehong Park ^{1,2*} , Seonmi Choi ¹ , Jun Takatoh ¹ , Shengli Zhao ³ , Andrew Harrahill ¹ , Bao-Xia Han ³ , Fan Wang ^{1*}
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15	The PDF file includes:
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1/ 10	Materials and Methods Figs. S1 to S6
10	$\frac{195.51}{References} (46-48)$
20	References (40-40)
21	Other Supplementary Materials for this manuscript include the following:
22	
23	MDAR Reproducibility Checklist
24	Movies S1 to S3
25	

26 Materials and Methods

27 Experimental models and subject details

- 28 All animal experiments were performed in accordance with the MIT Committee for Animal Care
- 29 Use Committee and Duke University Institutional Animal Care. Pups (postnatal 10 ~17 days) of
- 30 either C57BL/6 or tdTomato reporter mice (Ai14, Stock No: 007914, Jackson laboratory) were
- 31 used for tracing premotor neurons of the laryngeal muscles. Male homozygous Fos^{TVA} (Stock No:
- 32 027831, Jackson laboratory) were used for most of CANE experiments except for preRAm^{VOC}
- tracing. Male heterozygous Fos^{TVA} (crossed with C57BL/6 background) were used for preRAm^{VOC}
 tracing. Vglut2-ires-Cre mice (Stock No: 016963, Jackson laboratory) were crossed with Fos^{TVA}
- 35 mice to obtain Fos^{TVA} (het)/Vglut2-ires-Cre (het) for a subset of experiments.
- 35 mice to obtain Fos^{TVA} (het)/Vglut2-ires-Cre (het) for a subset of e
- 36

37 Viruses

- 38 AAV2retro-pENN.AAV.hSyn.Cre.WPRE.hGH (Addgene #105553)
- 39 AAV2retro-phSyn1(S)-FlpO-bGHpA (Addgene #51669)
- 40 AAV2/8-CAG-Flex-oG (Addgene #48332, Duke Viral Vector Core)
- 41 AAV2/8-CAG-Flex-TVA-mCherry (Addgene #74292, Duke Viral Vector Core)
- 42 AAV2/8-hSyn-Flex-TeLC-P2A-EYFP-WPRE (Addgene #135391)
- 43 AAV2/8-hSyn-DIO-EGFP (Addgene #50457)
- 44 AAV2/8-nEF-Con/Foff 2.0-ChRmine-oScarlet (Addgene #137161)
- 45 AAV2/8-nEF-Con/Fon-ChRmine-oScarlet (Addgene # 137159)
- 46 AAV2/8-nEF-Coff/Fon-ChRmine-oScarlet (Addgene # 137160)
- 47 AAV2/8-TRE3G-GFP-GFE3 (This study)
- 48 AAV2/8-TRE3G-EGFP (This study)
- 49 EnvA (M21)-RV-ΔG-GFP (30)
- 50 CANE (lenti)-hSyn-Cre (30)
- 51 CANE (lenti)-hSyn-DIO-tTA (This study)
- 52

53 Method details

54 Stereotaxic virus injection surgery

- 55 Mice were initially anesthetized by isoflurane (3%), then further maintained by isoflurane (1-2%)
- 56 until the surgeries ended. The heads of mice were fixed at a stereotaxic frame (Model 963, David
- 57 Kopf Instruments), and the body temperatures were maintained at 37°C with a heating pad. The
- virus solution was stereotaxically injected with a pulled-glass pipette (Drummond, 5-000-2005)
- 59 using an oil-hydraulic pump (MO-10, Narishige).
- 60

61 Stereotaxic coordinates

- Anterior-Posterior and Medial-Lateral coordinates are from the Bregma. Dorsal-Ventral
 coordinates are from the brain surface.
- 64 Nucleus ambiguous (NA): AP: -6.4 mm, ML: -1.2 mm, DV: -4.8 mm
- 65 RAm: AP: -5.8 mm, ML: 1.2 mm, DV: -5.4 mm (20° AP angle)
- 66 PAG: AP: -3.3 mm, ML: 0.6 mm, DV: -2.4 mm (30° AP angle)
- 67

68 Head-post and optic fiber implantation

- 69 In cases of the head-fixed or optogenetic experiments, mice were implanted with a head post
- 70 (custom made steel). For optogenetic manipulations, optic cannulas (200 µm core, 0.4NA, RWD

- 71 Life Science) were implanted. The implantations were performed right after the virus injections.
- 72 Dental cement (C&B Metabond) was applied to the skulls to secure the implantations.
- 73

74 Three-step monosynaptic tracing for premotor neurons of laryngeal muscles in adult mice

- 75 Laryngeal premotor neurons in adult mice were traced by the three-step monosynaptic rabies virus
- tracing as previously described (23). Briefly, mice pups were anesthetized by isoflurane (3% for
- 77 induction and 1.5% for maintenance). Midline incision in the neck skin and sternohyoid muscle
- 78 was performed, and the incised sternohyoid muscle was bilaterally retracted with thin thread to 79 expose the larynx. AAV2retro-hSyn-Cre was injected into laryngeal muscles (500 nl) using a
- a quartz micropipette (Sutter Instrument) through a micro syringe pump system (UMP3 and Micro4;
- 81 WPI). Three weeks or more after the AAV injection, a mixture of AAV2/8-CAG-Flex-oG and
- 82 AAV2/8-CAG-Flex-TVA-mCherry (120nl total with 1:1 ratio in volume) was stereotaxically
- 83 injected in the ipsilateral NA. Two weeks later, EnvA (M21)-RV- Δ G-GFP (200 nl) was injected
- 84 in the same injection target. After 5 days, the mice were perfused for histology.
- 85

86 Registering neurons in the Allen CCF

- 87 Registrations of laryngeal premotor and RAm^{VOC} neurons were performed as described previously
- 88 (23). Briefly, all neurons in serial-sectioned (80 μm) brain slices were manually registered to
 89 generate 3D coordinates in the Allen CCF with custom-written MATLAB. A Python package,
- 90 Brainrender2 (45) was used to visualize neurons in 3D.
- 91

92 Analysis of spatial distribution and correlation

- As previously described (23), a kernel density estimation in three-dimension was applied to the 3D-coordinates of registered cells. For 2D density plots, the 3D density estimations were projected to 2D dimension (AP, ML, or DV). The 3D density estimations were vectorized, then cosine similarities were calculated between each premotor map to plot a cross-correlogram. The coordinates of jaw and tongue premotor neurons were obtained from previous work.
- 98

99 Histology

- 100 Mice were anesthetized with an overdose of isoflurane and perfused with ice cold 1xPBS, followed
- 101 by 4% PFA. The brains were frozen in OCT compound (Sakura Finetek). Eighty-micron serial
- 102 coronal sections were made. Neurotrace blue (1:500, Thermo Fisher Scientific, N21479) was used
- 103 to visualize neuronal structures.
- 104

105 Immunohistochemistry (IHC) for ChAT and Fos

- 106 Free-floating IHC was performed as previously described (46). Coronal brain slices were 107 permeabilized for 3 hours in 1% Triton X in PBS (PBST), followed by the blocking solution (10% 108 Blocking One (Nacalai Tesque) in 0.3% PBST). Floating sections were incubated at 4 degrees for 109 24 hours with the primary antibody in the blocking solution, then washed with 1xPBS three times for 10 mins each. Secondary antibodies in the blocking solution were applied to the sections for 110 111 24 hours at 4 degrees. Tissue sections were rinsed with 1xPBS three times for 10min each. The 112 washed sections were mounted on slides with Mowiol. Antibodies for ChAT staining: primary 113 (Goat, 1:500, AB144P, Sigma) and secondary (anti-Goat, 1:500, Alexa FluorTM 555, A21432,
- 114 Invitrogen). Antibodies for Fos staining: primary (Rabbit, 1:4000, 2250S, cell signaling) and
- 115 secondary (anti-Rabbit, 1: 500, Alexa FluorTM Plus 647, A32795, Invitrogen).
- 116

117 Fluorescent HCR (v.3.0, Molecular Instruments) RNA-FISH

118 HCR was performed as previously described (46). In brief, floating brain sections were perfused 119 in 70% Ethanol/PBS overnight at 4 °C. The sections were washed with DEPC-PBS for three min 120 each. The sections were then treated with 5% SDS/DEPC-PBS for 45 min at room temperature. 121 After rinsing in $2 \times SSC$, the sections were incubated in $2 \times SSC$ for 15 min. The sections were then 122 incubated in probe hybridization buffer for 30 min at 37 °C for 30 min, followed by incubation 123 with probes (Fos, Vglut2, Vgat, Molecular Instruments) overnight at 37 °C. After washing in HCR 124 probe wash buffer (four times for 15 min at 37 °C), the sections were rinsed in 2× SSC (twice for 125 5 min) and incubated in HCR amplification buffer for 30 min at room temperature. The sections 126 were then incubated for 48 hours at 25 °C with appropriate hairpins conjugated with Alexa Fluor 127 (denatured and snap-cooled according to manufacturer's instructions) to visualize hybridization 128 signals. The washed sections with 2× SSC (twice) were mounted on slides with Mowiol.

129

130 Courtship male mice behaviors

131 Male mice were placed in a glass cylindrical chamber and acclimated for 10min before being

- 132 introduced to female partners. Female mice were placed in the chamber for up to 1 hour. The
- behaviors of the mice were recorded with a camera at 20 frames/s. Ninety-minutes or two hours
- 134 after introduction of female and the vocalization onsets, the male mice were perfused for *Fos* HCR 135 or Fos immunostaining, respectively.
- 135 136

137 CANE based targeting of RAm^{VOC}-neurons

- 138 Prior to CANE mediated capturing of RAm^{VOC}, each virgin male Fos^{TVA} mouse was first exposed 139 to a female mouse overnight and then isolated in a single chamber for one week to facilitate male
- 139 to a female mouse overnight and then isolated in a single chamber for one week to facilitate male 140 vocalization in the subsequent courtship contexts. Male mice were introduced with receptive
- females in a cylindrical chamber to elicit USVs for up to one hour. Two hours after the vocalization
- 142 onsets, CANE (lenti)-hSyn-Cre and Cre dependent AAV2/8-gene X (600 nl total with 4:1 ratio in
- volume; gene X: hSyn-Flex-TeLC-P2A-EYFP-WPRE, hSyn-DIO-EGFP, nEF-Con/Foff 2.0-
- 144 ChRmine-oScarlet) were stereotaxically injected to the RAm. For specifically targeting excitatory
- 145 Ram^{VOC} neurons, Fos^{TVA}/Vglut2-ires-cre mice were used, and CANE (lenti)-hSyn-DIO-tTA and
- 146 AAV2/8-TRE3G-geneX (600 nl total with 4:1 ratio in volume; gene X: GFP-GFE3, GFP), were
- 147 injected to the RAm.
- 148

149 **PreRAm**^{VOC} tracing

150 The procedure is the same as the other experiment using CANE to express helper viruses AAV2/8-

- 151 CAG-Flex-oG and AAV2/8-CAG-Flex-TVA-mCherry in RAm^{VOC}, followed by stereotaxic 152 injection of EnvA (M21)-RV- Δ G-GFP (200 nl) to RAm two weeks later.
- 153

154 Recording and analysis of USVs

- 155 USVs were recorded with a recording system for ultrasonic-range audio signals 156 (CM16/CMPA48AAF-5V, Avisoft-Bioacoustics). The audio signals were digitized at 250 kHz
- 157 with an analog-digital converter (PCIe-6321, National Instruments). Spectrogram of audio signals
- 158 were calculated by the Short Time Fourier Transform algorithm (512 Hanning window with 25%
- 159 overlap). USVs were detected by manual selection from the spectrograms within 30-125 kHz.
- 160 Classification of RAm^{VOC}-USVs were manually performed based on the criteria previously
- 161 described (34). Four acoustic features were calculated for each USV syllable: 1) loudness (average
- 162 band power between minimum and maximum frequency of each USV syllable as dB (relative to

background noise in the recording)), 2) spectral purity (relative power of dominant frequency), 3)

164 mean frequency (averaged dominant frequency at each time point), and 4) pitch variance (the

- 165 variance of dominant frequencies). Putative inspiratory vocalizations were manually selected,
- based on the two criteria: 1) time-locked to inspiration periods and 2) broad spectral representation.
- 167

168 **Respiratory activity recording and analysis**

169 Respiratory Activity was measured as previously described (5). Briefly, awake mice were head-170 fixed, and an airflow sensor (AMW330V, Honeywell) was closely positioned to the nose of the 171 mice. Voltage signals from the sensor were recorded at 250 kHz (PCIe-6321, National 172 Instruments) and down-sampled to 1kHz for analysis. All breathing signals were normalized by 173 their resting states: the breathing signals were subtracted by the reference value (at no-flow) and 174 divided by the standard deviation of the resting breathing. For labeling flat-expirations, custom 175 Julia codes were used to automatically detect flatten respiratory periods. Each negative and 176 positive period of the breathing signals was interpolated and labeled as inspiration ($-\pi$ to 0) and 177 expiration (0 to π) phases, respectively. Inspiration peaks were defined as the minimum values 178 during each inspiration period. The inspiration peaks were interpolated to visualize the amplitude 179 changes over time in average.

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181 Correlation between duration of USVs and expirations

- 182 A linear regression model was used to fit a model of duration of USVs and flat-expirations. R^2 was
- 183 calculated to assess the model.
- 184

185 Calculation of laser stimulation phases

186 Laser stimulation phases with respect to respiration (Φ_{laser}) were similarly calculated as previously

- 187 described (18). Briefly, each negative and positive period of the breathing signals was interpolated 188 and labeled as inspiration ($-\pi$ to 0) and expiration (0 to π) phases, respectively. Laser stimulation
- time relative to the onset of the inspiration was projected on the prior (control) respiratory period
- 190 to define Φ_{laser} as from $-\pi$ to π . Each latency and duration of RAm^{VOC}-USVs and expiration data

191 with respect to the laser stimulation phases was polynomial fitted using CurveFit.jl package to

- 192 visualize the curves of the data.
- 193

194 **Respiratory phase maps of USVs**

195 The onset and offset time of USV syllables were projected onto the respiratory phase map. 196 Vocalizations are classified as inspiratory or expiratory vocalization based on the phase values

- 196 Vocalizations are classified as inspiratory or expiratory vocalization based on t197 (negative as inspiratory and positive as expiratory)
- 198

199 Pain-induced audible squeak experiments

200 Either tail-pinch or electrical foot shock were applied to the mice. For tail-pinch experiments, 201 awake mice were head-fixed and allowed to run on a running wheel. Mice tails were gently 202 grabbed with a globed-hand and further pinched to elicit squeaks. Respiratory activities of the mice 203 were measured with the airflow sensor. For electrical foot shock experiments, mice were placed 204 in a foot-shock chamber, and brief electrical foot shock were delivered to the mice ($\leq 2s, 0.5mA$). 205 The behaviors of the mice in the chamber were recorded with a camera (with audible mic) at 20 206 frames/s. The squeaks from both stimuli were audible and also represented in the USV spectrum 207 range.

207 r 208

209 **Abdominal EMG recordings**

210 Mice were initially anesthetized by isoflurane (3%), then further maintained by intraperitoneal 211 injection of the ketamine and xylazine mixture (1 and 0.1 mg/kg, respectively). The skin above 212 abdominal muscles were shaved and opened to expose abdominal muscles. Teflon coated silver 213 wires (bare diameter: 76.2 µm, AM systems cat. 785500) were used to record EMG. The insulation 214 was removed from the tips of silver wires (2mm) for recording. Recording wire was inserted into 215 the abdominal muscle, while reference wire was inserted between the skin and fascia above the 216 muscle. AC Amplifier (DAM80, World Precision Instruments) was used to record EMG, and the 217 voltages were filtered (high pass: 100Hz, low pass: 10kHz) and collected with the same DAQ 218 board (PCIe-6321, National Instruments). The sampling rate for EMG was 250kHz for 219 simultaneous recording of USVs. The voltage recordings were down-sampled to 20kHz for 220 analysis. The root-mean-square filter was applied to visualize the EMG responses. Averaged EMG 221 responses during PAG stimulation (2s) were normalized by averaged resting EMG responses (1s) 222 to calculated PAG-evoked EMG.

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224 Vocal cord imaging and analysis

225 Mice were initially anesthetized by isoflurane (3%), then further maintained by intraperitoneal 226 injection of the ketamine and xylazine mixture (1 and 0.1 mg/kg, respectively). The heads of mice 227 were fixed with clamps, and the mice were put on a flat platform. A round post was placed under 228 the neck to keep the axis of the oral cavity and trachea straight. The tongue was gently pulled out 229 and moved down with a flat metal depressor (custom made) to help visualize the vocal cords. An 230 optic fiber was attached to the tip of the depressor to illuminate the inside of the oral cavities with 231 a red LED (635 nm, Doric). A camera (acA640-750um, Basler) with a lens (Basler Lens, C23-232 3520-2M-S f50mm) was used to image the vocal cords. Vocal cords were imaged at 100 233 frames/sec. The glottal areas of the vocal cords were calculated by tracking the videos using 234 DeepLabCut (47).

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Optogenetic stimulation of RAm^{VOC} and PAG^{RAm/Vglut2} 236

Awake mice were head-fixed on a running wheel, and respiratory activities and sound productions 237 were measured together. Bilateral (RAm^{VOC}-ChRmine) optogenetic stimulation was applied 238 239 through optic fibers (0.39 NA, 200um core). 560 nm laser (less than 10mW at the tips) was used, 240 and the stimulation parameters were modulated by TTL pulses with PulsePals. In experiments with 241 RAm^{VOC}-GFE3 or control mice, optogenetic stimulation of the PAG was used to elicit USVs in a 242 head-fixed setup. AAV2retro-hSyn-FlpO was injected into the RAm, and Cre/Flp-codependent 243 AAV2/8-nEF-Con/Fon-ChRmine-mScarlet was injected into the PAG. 244

245 **Statistics**

- 246 All data are represented in mean±s.e.m. Statistical analyses were performed in Julia using HypothesisTests.jl package. Non-parametric test, Mann-Whitney was used to compare respiratory 247 modulation in RAm^{VOC}-TeLC and RAm^{VOC}-GFP mice; mean frequency of RAm^{VOC}-USVs over 248 female-directed USVs; and changes in inspiration peaks in RAm^{VOC}-GFE3 and RAm^{VOC}-GFP 249 250 mice.
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Fig. S1. Distribution of laryngeal premotor neurons and their collaterals to the other orofacial motor nuclei.

259 (A) 3D reconstructed distribution of laryngeal premotor neurons (magenta, aqua, and gold for 3 260 different mice) reconstructed in the CCF. (B) Spatial density distributions of the laryngeal 261 premotor neurons (red) together with masseter (green) and genioglossus (blue) premotor neurons. 262 (C) Cross-correlation of 3D spatial distributions of three orofacial premotor neurons (three 263 examples for each: L (Larynx), M (Masseter), and G (Genioglossus). (D) Axon collaterals of rabies-labeled laryngeal premotor neurons (green) in other orofacial motor nuclei. The motor 264 nuclei were revealed with ChAT immunolabeling (red). Neurotrace Blue was used to visualize 265 neuronal structures. All scale bars indicate 200 µm. 266



269 Fig. S2. Vocalization induced strong *Fos*+ expression in the RAm but not in the preBötC.

Fluorescent in situ hybridization of *Fos* (green) in the preBötC (dash-circled, upper left) and RAm (bottom left) after female-directed male USVs. The right columns show *Fos* expression in the same regions without vocalization behaviors (male mice in a chamber alone). Note that no discernable *Fos* expression was found in the preBötC. Neurotrace Blue (magenta) was used to visualize

- 274 neuronal structures. All scale bars indicate 200 μm.



Fig. S3. RAm^{VOC}-TeLC mice were still able to modulate respiratory activity during running.

(A) Schematic for encouraging mice to run for respiratory modulation. (B) Respiratory recordings
in RAm^{VOC}-GFP (upper) and RAm^{VOC}-TeLC (bottom) mice. Magenta lines indicate the onsets of
tail-grabbing. Note that downward flows correspond to inspirations. (C) Changes in inspiration
(left) and expiration (middle) amplitude, and frequency (right) of respiratory flows. RAm^{VOC}TeLC mice (green, n=3) and RAm^{VOC}-GFP control mice (grey, n=4)



Fig. S4. PAG stimulation-evoked USVs and abdominal muscle activity were both abolished in RAm^{VOC}-TeLC mice.

(A) RAm^{VOC} axon terminals (GFP+) were observed in the thoracic spinal segment near motoneurons (ChAT+). (B) Schematic for eliciting USV in mice via PAG opto-stimulation. To target RAm projecting PAG neurons for vocalization, retrograde AAV-Flp and Flp dependent AAV- ChRmine were injected into the PAG and RAm, respectively. For RAm^{VOC} targeting, CANE-Cre and AAV-Flex-TeLC or GFP were injected into the RAm to express TeLC or GFP in RAm^{VOC}-neurons. (C) Sound and abdominal EMG recording during 2s PAG stimulation in anesthetized mice. (D) Spectrogram (upper) and EMG responses (RMS filtered, bottom) of GFP control and TeLC mice during PAG stimulation (2s, lime shades). (E) Evoked EMG responses (voltage) during PAG stimulation. RAm^{VOC}-TeLC mice (green, n=3) and RAm^{VOC}-GFP control mice (grey, n=3) Averaged EMG responses during PAG stimulation (2s) were normalized by averaged resting EMG responses (1s) to calculated PAG-evoked EMG.

Rabies labeled laryngeal premotor neurons in preBötC (RV-GFP) are Vgat+



344 Fig. S5. preBötC contains *Vgat*+ inhibitory laryngeal premotor neurons.

345 Green neurons are laryngeal premotor neurons labeled with monosynaptic rabies-GFP through

346 transsynaptic tracing (as described in Figure 1), and the in-situ hybridization signal with *Vgat*

- 347 probe is shown in magenta.



355 Fig. S6. Spontaneous vocalizations emerged in the RAm^{VOC}-GFE3 mice.

(A) Schematic for recording male mice vocalization in an isolation chamber. (B) Raw audio data
(upper) and corresponding spectrogram plots (bottom). (C) An example of a single USV syllable
from the spontaneous vocalizations. (D) Rates of spontaneous vocalizations in RAm^{VOC}-GFE3
mice (n=6) and RAm^{VOC}-GFP mice (n=4). Note that no spontaneous vocalization was found in
the GFP control mice. (E) Durations of the spontaneous vocalizations of six RAm^{VOC}-GFE3

- 361 mice. Single dots show outliers.

Movie S1. A RAm^{VOC}-GFP mouse in a foot-shock chamber. Raw audio signals (upper left) and corresponding spectrogram (bottom left, in ultrasonic range). Sounds in the video (right)

- 366 were recorded with an audible microphone.
- Movie S2. A RAm^{VOC}-TeLC mouse in a foot-shock chamber. The mouse showed behavioral
 responses to foot-shock but did not produce audible squeaks.
- 369 Movie S3. Glottal responses to RAm^{VOC}-ChRmine stimulation. Four red dots are used to 370 trace the glottal area. Red lines represent the glottal area. Green bars indicate the optogenetic
- 371 stimulation periods.

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