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### 30 Abstract

31 Obstructive sleep apnea (OSA) is characterized by sporadic collapse of the upper airway 32 leading to periodic disruptions in breathing. Upper airway patency governed by genioglossal 33 nerve activity originates from the hypoglossal motor nucleus. Mice with targeted deletion of the 34 gene Hmox2, encoding the carbon monoxide (CO) producing enzyme, heme oxygenase-2 (HO-35 2), exhibit severe OSA, yet the contribution of central HO-2 dysregulation to the phenomenon is 36 unknown. Using the rhythmic brainstem slice preparation, which contains the preBötzinger 37 complex (preBötC) and the hypoglossal nucleus, we tested the hypothesis that central HO-2 38 dysregulation weakens hypoglossal motoneuron output. Disrupting HO-2 activity increased 39 transmission failure as determined by the intermittent inability of the preBötC rhythm to trigger 40 output from the hypoglossal nucleus. Failed transmission was associated with a reduced input-41 output relationship between the preBötC and the motor nucleus. These network phenomena 42 were related to smaller inspiratory drive currents and reduced intrinsic excitability among 43 hypoglossal neurons. In addition to HO-2, hypoglossal neurons also expressed the CO-44 regulated  $H_2S$  producing enzyme cystathionine  $\Box$ -lyase (CSE).  $H_2S$  abundance was higher in 45 hypoglossal neurons of HO-2 null mice than wild-type controls. Disrupting CSE function 46 normalized transmission in HO-2 null mice and an H<sub>2</sub>S donor mimicked the effects of HO-2 47 dysregulation. These findings demonstrate a hitherto uncharacterized modulation of hypoglossal activity through the interaction of HO-2 and CSE-derived  $H_2S$ , and supports the 48 49 perspective that centrally derived HO-2 activity plays an important role regulating upper airway 50 control.

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### 56 Introduction

57 Obstructive sleep apnea (OSA) is a prevalent breathing disorder affecting nearly a billion 58 people throughout the world<sup>1,2</sup>. It is characterized by the periodic cessation of breathing during 59 sleep due to sporadic upper airway collapse. When left untreated, OSA predisposes the 60 individual to a variety of diseases including hypertension<sup>3,4</sup>, diabetes<sup>5,6</sup>, and cognitive decline<sup>7,8</sup>. 61 Multiple factors contribute to the genesis of OSA including compromised pharyngeal anatomy 62 <sup>9,10</sup>, inadequate upper airway muscle function<sup>11-13</sup>, low arousal threshold<sup>14</sup>, and a hypersensitive 63 chemoreflex (i.e., high loop gain <sup>15</sup>).

64 Peng et al. recently reported that mice with deletion of the Hmox 2 gene, which encodes the enzyme heme oxygenase 2 (HO-2), exhibit a high incidence of OSA<sup>16</sup>. OSA in HO-2 null 65 66 mice was attributed in part to the increased loop gain arising from the heightened carotid body chemo reflex<sup>16-19</sup>. While HO-2 produces several bioactive products<sup>20</sup>, the enhanced carotid 67 body chemo reflex and subsequent OSA phenotype is attributed to the loss of HO-2 dependent 68 CO production<sup>16</sup>. However, we hypothesize that additional factors, such as changed activities in 69 70 the central nervous system, may also contribute to the OSA phenotype observed in HO-2 null 71 mice.

Sporadic loss of neuromuscular control over upper airway muscles is a key contributor to producing obstructive apneas<sup>21,22</sup>. Predisposition to intermittent reductions in airway patency can originate from changes in CNS activity. Disrupting rhythmic excitation of the hypoglossal nucleus that drives genioglossal nerve activity increases the likelihood for the tongue to occlude the upper airway during inspiration. Such disruptions may involve changing the statedependent balance between excitation and inhibition received by respiratory hypoglossal motor

neurons<sup>23</sup> and/or the direct modulation of intrinsic excitability of hypoglossal neurons<sup>24-26</sup>. It,
however, is unknown whether impaired HO-2 signaling in the hypoglossal nucleus influences
synaptic and/or intrinsic neuronal properties to alter output from the motor nucleus that may
ultimately contribute to upper airway obstruction.

82 We tested this possibility using a combination of electrophysiological, genetic, and 83 pharmacological approaches in the rhythmic medullary brainstem slice preparation. We found 84 that dysregulated HO-2 activity in the hypoglossal nucleus acts through CSE-dependent  $H_2S$ 85 signaling to reduce motor neuron excitability. This in turn, diminishes the input-output 86 relationship between the preBötC and hypoglossal nucleus, and increases the likelihood of 87 transmission failure between the premotor rhythm and motor nucleus output. These 88 observations indicate that hypoglossal HO-2 / CO and CSE / H<sub>2</sub>S activities interact as important 89 modulators of hypoglossal output that potentially contributes to changed upper airway tone 90 when dysregulated.

#### 91 Methods

*Study Approval.* In accordance with National Institutes of Health guidelines, all animal
protocols were performed with the approval of the Institute of Animal Care and Use Committee
at The University of Chicago (ACUP 72486, ACUP 71811).

95 **Experimental Animals.** Experiments were performed using neonatal (postnatal day 6 to 96 postnatal day 12) wildtype mice (C57/BL6; Charles River), HO-2 null mice (from S. H. Snyder; 97 The Johns Hopkins University), and HO-2:CSE double-null mice. HO-2:CSE double-null mice 98 were created by crossing HO-2 null females with CSE null males (from R. Wang, Department of 99 Biology, Laurentian University, Sudbury, ON, Canada). Tissues from both sexes were used. No 100 sex-based differences were observed; therefore, all sexes were pooled for analysis. All litters 101 were housed with their dam in ALAAC-approved facilities on a 12 hour / 12-hour light-dark 102 cycle.

103 Pharmacological Agents. Heme oxygenase activity was blocked using bath application of 104 Chromium (III) Mesoporphyrin IX chloride (ChrMP459, 20uM; Frontiers Sciences, Newark DE). 105 CORM-3 (20µM; Sigma-Aldrich St. Louis MO), a CO-donor, was bath applied following 106 ChrMP459 application. NaHS (10 $\mu$ M to 100  $\mu$ M; Sigma-Aldrich), a H<sub>2</sub>S donor, was bath 107 applied. In all patch clamp experiments, fast synaptic glycinergic and GABAergic inhibition was 108 blocked by bath application of strychnine (1µM; Sigma-Aldrich), and picrotoxin (50µM; Sigma-109 Aldrich), respectively, Inhibition of CSE production was accomplished by in vivo L-110 propargylglycine (L-PAG, 30mg/kg (Sigma-Aldrich) administered (*i.p.* injection) 2.5 to 3 hrs prior

to preparation of the rhythmic brainstem slice preparation. Inhibition of potassium channels  $SK_{Ca}$  and ATP-sensitive potassium channel ( $K_{ATP}$ ) was via bath application of APAMIN (200  $\mu$ M; Sigma-Aldrich) and Tolbutamide (100  $\mu$ M; Sigma-Aldrich) respectively.

114 *Measurement of H<sub>2</sub>S Production.* Coronal brainstem sections (300 $\mu$ m thick) were cut with a 115 cryostat at -20°C. The hypoglossal nucleus and control brainstem regions were excised with a 116 chilled micro-punch needle. Hypoglossal tissue from a single brainstem was not sufficient for 117 effectively measuring H<sub>2</sub>S levels; therefore, we pooled micro punched tissue from two mice for 118 each sample where  $H_2S$  levels measured. H<sub>2</sub>S levels were determined as described 119 previously<sup>27</sup>. Briefly, cell homogenates were prepared in 100 mM potassium phosphate buffer 120 (pH 7.4). The enzyme reaction was carried out in sealed tubes. The assay mixture in a total 121 volume of 500µL contained (in final concentration): 100 □ mM potassium phosphate buffer (pH 122 7.4), 800µM□L-cysteine, 80µM pyridoxal 5'-phosphate with or without L-PAG (20µM) and cell 123 homogenate (20µg of protein), was incubated at 37°C for 1 □hr. At the end of the reaction, 124 alkaline zinc acetate (1% mass / volume; 250µL) and trichloroacetic acid (10% vol/vol) were 125 sequentially added to trap  $H_2S$  and stop the reaction, respectively. The zinc sulfide formed was 126 reacted with acidic N,N-dimethyl-p-phenylenediamine sulfate (20µM) and ferric chloride (30µM) 127 and the absorbance was measured at 670 nm using Shimadzu UV-VIS Spectrophotometer. L-128 PAG inhibitable H<sub>2</sub>S concentration was calculated from a standard curve and values are 129 expressed as nanomoles of H<sub>2</sub>S formed per hour per mg of protein.

**Immunohistochemistry.** Anaesthetized mice (urethane, 1.2g•kg<sup>-1</sup> *i.p.*) were perfused transcardially with heparinized phosphate-buffered saline (PBS) for 20 min followed by 4% paraformaldehyde in PBS. Brainstems were harvested, post-fixed in 4% paraformaldehyde overnight, and cryoprotected in 30% sucrose/PBS at 4°C. Frozen tissues were serially sectioned at a thickness of 20µm (coronal section) and stored at –80°C. Sections were treated with 20% normal goat serum, 0.1% bovine serum albumin and 0.1% Triton X-100 in PBS for 30

min and incubated with primary antibodies against choline acetyltransferase (ChAT, 1:100;
Millipore; #AB144P), HO-2 (1:200, Novus Biologicals; # NBP1-52849) and CSE (1:250; gift
from Dr. Schenider) followed by Texas Red-conjugated goat anti-mouse IgG or FITCconjugated goat anti-rabbit IgG (1:250; Molecular Probes). After rinsing with PBS, sections were
mounted in Vecta shield containing DAPI (Vector Labs) and analyzed using a fluorescent
microscope (Eclipse E600; Nikon).

142 Brainstem Slice for Electrophysiology. The isolated rhythmic brainstem slice was prepared as previously described <sup>28</sup>. Briefly, animals were euthanized by decapitation. Brainstems were 143 144 rapidly dissected, isolated, and placed into ice cold artificial cerebral spinal fluid (aCSF) 145 (composition in mM: 118 NaCl, 25 NaHCO<sub>3</sub>, 1 NaH<sub>2</sub>PO<sub>4</sub>, 1 MgCl<sub>2</sub>, 3 KCl, 30 Glucose, 1.5 CaCl<sub>2</sub>, 146 pH=7.4) equilibrated with 95%  $O_2$ , 5%  $CO_2$ . The isolated brainstem was glued to an agar block (dorsal face to agar) with the rostral face up and submerged in aCSF equilibrated with 147 148 carbogen. Serial cuts were made through the brainstem until the appearance of anatomical 149 landmarks such as the narrowing of the fourth ventricle and the hypoglossal axons. The 150 preBötC and XIIn was retained in a single transverse brainstem slice (thickness:  $560 \pm 40 \mu m$ ). 151 The slice was transferred into the recording chamber (~6mL volume) where it was continuously 152 superfused with recirculating aCSF (flow rate: 12 -15mL/min). Prior to recording, extracellular 153 KCI was raised to 8mM and the spontaneous rhythm was allowed to stabilize prior to the start of 154 every experiment.

**Electrophysiology.** Extracellular population activity was recorded with glass suction pipettes filled with aCSF. Pipettes were positioned over the ventral respiratory column containing the preBötC and over the medial dorsal column containing the hypoglossal nucleus. In some experiments, a third pipette was positioned between the preBötC and hypoglossal nucleus just lateral to the axon tract to record transmission through the premotor field containing intermediate premotor inspiratory neurons<sup>29,30</sup>. Extracellular population activity was recorded

with glass suction pipettes filled with aCSF<sup>31</sup>. The recorded signal was sampled at 5kHz,
amplified 10,000X, with a lowpass filter of 10 kHz using an A-M instruments (A-M Systems,
Sequim, WA, USA) extracellular amplifier. The signal was then rectified and integrated using
Clampfit electronic filter. Recordings were stored on a computer for *posthoc* analysis.

165 All intracellular recordings were made using the Multiclamp 700B (Molecular Devices: 166 https://www.moleculardevices.com/systems/conventional-patch-clamp/multiclamp-700b-

- <u>microelectrode-amplifier</u>). Acquisition and post hoc analyses were performed using the Axon
   pCLAMP10 software suite (Molecular Devices: https: <u>www.moleculardevices.com/system/axon-</u>
   conventional-patch-clamp/pclamp-11-software-suite).
- 170 Whole cell patch clamp recordings of hypoglossal motor neurons were obtained using the blind-171 patch technique with a sample frequency of 40 kHz. Recordings were made with unpolished 172 patch electrodes, pulled from borosilicated glass pipettes with a capillary filament<sup>31</sup>. The 173 electrodes had a resistance of 3.5-8 M $\Omega$  when filled with the whole cell patch clamp pipette 174 solution containing (in mM): 140 K-gluc acid, 1 CaCl<sub>2</sub>, 10 EGTA, 2 MgCl<sub>2</sub>, 4 Na<sub>2</sub>-ATP, 10 175 HEPES. Patch clamp experiments were performed with a patch clamp amplifier (Multiclamp 176 700B, Molecular Devices, Sunnyvale, CA, USA), and the software program pCLAMP 10.0 177 (Molecular Devices). Neurons located at least two to three cell layers (about 75-250µm) rostral 178 from the caudal surface of the slice were recorded. The liquid junction potential was calculated 179 to be -12mV and was subtracted from the membrane potential. The series resistance was 180 compensated and corrected throughout each experiment. In voltage clamp experiments, 181 membrane potential was held at -60mV. Current clamp experiments used a holding potential 182 between 0 and -100pA to establish the baseline resting membrane potential between -55 and -183 70mV. In some cases, we determined rheobases using a ramp protocol in our current clamp 184 recordings. This ramp protocol consisted of a hyperpolarizing step (-100pA) succeeded by the 185 injection of a ramping depolarizing current (122pA/sec; peak current 600pA).

186 Statistical Analyses. Unless otherwise explicitly stated elsewhere, each n value represents an 187 individual animal that served as a biological replicate for a given measurement. Transmission 188 was expressed as a percentage of the hypoglossal network bursts corresponding to the total 189 network bursts from either the preBötC or the premotor field. Burst were considered 190 corresponding if initial start time of bursts were within 500-750ms of each other (corresponding 191 time was maximized until only one hypoglossal burst per preBötC was detected). Mean I/O and 192 transmission values for each slice were calculated using a 120 s window. This window was 193 taken at the end of each baseline or pharmacological agent phase (each phase duration = 194 600 sec). The input-output (I/O) ratio for each inspiratory event (defined by network a burst in preBötC) was calculated as previously described in <sup>31</sup>. I/O ratios values for preBötC bursts 195 196 without a corresponding XIIn burst were designated as 0. To illustrate the cycle-to-cycle input-197 output relationships between networks, heat maps of I/O ratio values were plotted for each slice 198 included in the experiment. Each row represents sequential cycles from a single slice 199 experiment. As the rhythmic frequency across preparations varied, the number of events (i.e., 200 cycle number) in the 120 s analysis window also varied; therefore, either the total number of 201 cycles or 25 consecutive cycles from in a given analysis window were plotted.

202 Statistics were performed using Origin 8 Pro (OriginLab, RRID:SCR 014212) or Prism 6 203 (GraphPad Software; RRID:SCR 015807). In cases where the distribution of data appeared 204 normal, comparisons between two groups were conducted using either paired or unpaired two-205 tailed t-tests as appropriate. In cases, where the distribution of individual data points did not 206 appear normal, the Wilcoxon match-paired signed rank test was performed. A one-way 207 ANOVA was performed followed by *posthoc* Dunnett's test comparing experimental groups to 208 control when a comparison of three or more groups. In plots where the mean ± S.E.M. are 209 presented, the mean and S.E.M. are overlaid on the individual results from the corresponding 210 dataset. Differences were defined when the P-value was less than 0.05.

### 211 Results

Hypoglossal neurons express hemeoxygenase-2 (HO-2). We first assessed whether hypoglossal neurons express HO-2. Hypoglossal neurons showed positive immunohistochemical expression for HO-2 as indicated by co-localization of HO-2 with Choline acetyl transferase (ChAT), an established marker of these neurons <sup>11</sup> (**Fig.1A**, n=3).

216 **Disrupting HO-2 function impairs hypoglossal inspiratory activity.** Extracellular field 217 recordings in the rhythmic brainstem slice were simultaneously recorded from the preBötC and 218 the corresponding motor output from the hypoglossal nucleus. Two approaches were employed 219 to assess the role of HO-2: (1) using Cr(III) Mesoporphyrin IX chloride (ChrMP459, 20µM), an 220 inhibitor of HO<sup>32</sup>; and (2) using brain slices from HO-2 null mice.

221 Representative extracellular field recordings from the preBötC and hypoglossal nucleus prior 222 and during ChrMP459 exposure were shown in Fig 1B (n=11). While ChrMP459 suppressed 223 hypoglossal burst amplitude (Fig 1C, left, n=11; Baseline: 99.57 ± 0.60%, ChrM459: 81.80 ± 224 10.80%, P=0.007), the HO inhibitor had no effect on preBötC burst amplitude (Fig 1C, right, 225 n=11; Baseline: 99.93 ± 0.34%, ChrMP459: 93.23 ± 4.80%, P=0.21). ChrMP459 consistently 226 reduced the cycle-to-cycle input-output relationship between preBötC and the motor nucleus as 227 revealed by examining the cycle-to-cycle I/O across preparations (Fig 1D, Baseline:  $1.00 \pm 0.07$ 228 vs. ChrMP459: 0.55 ± 0.11; P=.006). In the extreme, altered input-output relationships between 229 preBötC and the hypoglossal nucleus may increase the propensity for transmission failures as 230 determined by the inability of preBötC activity to produce hypoglossal output at the network level<sup>31</sup>. Indeed, the reduced I/O ratio was associated with an increase in failed transmissions of 231

the preBötC activity to output from the hypoglossal nucleus (**Fig 1E**, Baseline:  $90.85 \pm 5.80\%$ vs. ChrMP459:  $80.21 \pm 7.40\%$ , P=0.039). Together these findings suggest that HO inhibition causes a generalized weakening in the activity relationship between the rhythm generating network and hypoglossal motoneurons.

236 ChrMP459 is a pan HO inhibitor; however, it cannot distinguish activities between heme 237 oxygenase isoforms. To assess the specific contribution of HO-2, we compared rhythmic 238 activities in brain slices from wild type (n=9) and HO-2 null (n=7) mice (Fig 2A). Larger cycle-to-239 cycle I/O ratios were observed in wild type slices as compared to HO-2 null slices (Fig 2B: wild 240 type: 0.99 ± 0.04 vs. HO-2 null: 0.76 ± 0.11, P=0.045). Similarly, transmission of preBötC 241 activity to the hypoglossal nucleus was greater in wild type than in HO-2 null slices (Fig 2 C, 242 wild type: 96.53 ± 1.76% vs. HO-2 null: 62.55 ± 7.93%, P=0.0006). These findings established 243 that genetic elimination of HO-2 produces a similar phenomenon to that of pan HO inhibition 244 indicating that loss of HO-2 activity alone is sufficient for impairing transmission from the 245 preBötC to the hypoglossal nucleus. Given these similarities and the limited availability of HO-2 246 null mice, several of the following studies were performed using the ChrMP459 in rhythmic wild 247 type brainstem slices.

248 HO inhibitor does not affect premotor neuron activity. Intermediary premotor neurons relay drive from the preBötC to the hypoglossal nucleus<sup>29,30</sup>. Therefore, it was possible that HO 249 250 inhibition impaired transmission of drive from the preBötC by perturbing activity from 251 intermediary premotor neurons. To address this possibility, triple extracellular recordings (n=5) 252 were made from the preBötC, the field of the ipsilateral premotor neurons, and the hypoglossal 253 nucleus. Baseline transmission from the preBötC to the premotor field and to the hypoglossal 254 nucleus was reliable and consistent (Fig. 3A, middle panel). However, ChrMP459 disrupted 255 activity in the hypoglossal nucleus despite unaltered activities in either the preBötC or the 256 intermediate premotor field (Fig 3A, right panel). Indeed, while neither transmission failures nor

257 the cycle-to-cycle I/O ratio from the preBötC to the premotor field was affected by ChrMP459 258 (**Fig 3B**: *left*; I/O: Baseline:  $1.16 \pm 0.09$  vs ChrMP  $1.16 \pm 0.14$ , P=0.31; *right*; Transmission: 259 Baseline:  $100.0 \pm 0.0\%$  vs ChrMP 86.35 ± 11.81%, P=0.312), the HO inhibitor reduced the 260 transmission of activity and the cycle-to-cycle I/O ratio between the premotor field and the 261 hypoglossal nucleus (Fig 3C: left, I/O: Baseline: 1.12±.17 vs ChrMP 0.40±0.09, P=0.04; right, 262 Transmission: Baseline 89.57±1.3% vs ChrMP 57.62±13.69%, P=0.04). These results 263 suggested that dysregulated HO-2 modulates is a postsynaptic phenomenon in the hypoglossal 264 nucleus.

265 HO inhibition suppresses inspiratory drive currents and reduces excitability in 266 hypoglossal neurons. To examine the effect of HO inhibition on postsynaptic activity of the 267 hypoglossal neurons, we performed patch clamp recordings from a total of 27 wild type 268 hypoglossal neurons exposed to ChrMP459. These hypoglossal neurons were disinhibited from 269 fast inhibition (50µM picrotoxin and 1µM strychnine) allowing us to focus on inspiratory-related 270 fast glutamatergic drive. 19 of the 27 hypoglossal neurons received excitatory synaptic drive in-271 phase with the preBötC (i.e., inspiratory hypoglossal neurons). Peak inspiratory drive currents 272 were reduced in ChrMP459 (Fig 4A, n=19, Baseline: -142.9 ± 22.82 pA vs. ChrMP459: -95.31 ± 273 21.79 pA, P=0.004). Reduced drive coincided with hypoglossal neurons generating fewer 274 action potentials per preBötC burst in ChrMP459 (Fig 4B, n=17, Baseline: 14.68 ±2.23 action 275 potentials per burst vs. ChrMP459: 6.79± 1.54 action potentials per burst, P<0.0001). 276 Furthermore, as determined by the injection of a depolarizing ramp current into hypoglossal 277 neurons, the HO inhibitor increased rheobase among inspiratory hypoglossal neurons (Fig 4C, 278 n=19, Baseline: 167.5 ± 35.83 pA vs. ChrMP459: 338.0 ± 82.50 pA; P=0.008) yet decreased 279 rheobase in non-inspiratory hypoglossal neurons (i.e., neurons not receiving drive during 280 preBötC activity; Supplemental Fig 1, n=8, Baseline: 280.5±56.43 pA vs 228.2±47.96pA, 281 P=0.017).

282 Elevated levels of  $H_2S$  are observed in the hypoglossal nucleus of HO-2 null mice. We 283 next sought to determine the mechanism(s) by which inhibition of HO-2 affect hypoglossal neuron activity. Earlier studies<sup>33,34</sup> have reported that HO-2 is a negative regulator of CSE-284 dependent H2S production. To test this possibility, we first examined whether the hypoglossal 285 286 neurons express CSE. Brainstem sections from the wild type hypoglossal nucleus revealed 287 CSE hypoglossal tissue punches from wild type and HO-2 null mice. Relative to the wild type 288 hypoglossal nucleus, H<sub>2</sub>S is expressed in ChAT-positive hypoglossal neurons (Fig 5A, n=3). 289 We then determined H<sub>2</sub>S abundance in hypoglossal tissue punches from wild type and HO-2 290 null mice. Relative to the wild type hypoglossal nucleus (Fig 5B blue; n=6, 60.58 ± 6.37 nmol • mg<sup>-1</sup> • h<sup>-1</sup>), H<sub>2</sub>S abundance was greater in the hypoglossal nucleus of HO-2 null 291 292 mice (**Fig 5B** red; n=6, 144.12  $\pm$  8.29 nmol  $\cdot$  mg<sup>-1</sup>  $\cdot$  h<sup>-1</sup>), but not different from the inferior olive 293 brainstem region of HO-2 null mice (**Fig 5B** grey; n=4, 56.10  $\pm$  2.88 nmol  $\bullet$  mg<sup>-1</sup>  $\bullet$  h<sup>-1</sup>). These 294 findings suggest that the hypoglossal nucleus expresses CSE and HO-2 negatively regulates 295 H<sub>2</sub>S production in the hypoglossal nucleus.

# 296 $H_2$ S mediates impaired transmission of inspiratory drive caused by disrupted HO-2 297 function.

If the impaired transmission of inspiratory drive to the hypoglossal nucleus by disrupted HO-2 function involves CSE-derived H<sub>2</sub>S then: 1) a H<sub>2</sub>S donor should mimic the effects of disrupted HO-2 activity; 2) CO administration should improve the input-output relationship in respiratory slices from HO-2 null mice and ChrM459 application; and 3) CSE blockade should restore the transmission from the preBötC to the hypoglossal nucleus. The following experiments tested these possibilities.

Wild type brainstem slices exhibited a nearly 1:1 ratio of transmission of inspiratory activity from preBötC to hypoglossal neurons (**Fig 5C**, *left*). Application of NaHS, a H<sub>2</sub>S donor reduced transmission from preBötC to hypoglossal (**Fig. 5C**, *middle*, *right*) in a dose-dependent manner (**Fig. 5D**; 0 $\mu$ M NaHS: n=9, 100 ± 0.73%; 10 $\mu$ M NaHS: n=5, 90.14 ± 6.08%; 50 $\mu$ M NaHS: n=6,

308 84.0  $\pm$  3.91%; 100µM NaHS: n=9, 81.2  $\pm$  5.83%), which coincided with a reduction in I/O ratio 309 by NaHS (**Fig. 5E**; 0 µM NaHS: 1.055  $\pm$  0.028; 10µM NaHS: 0.85  $\pm$  0.09; 50µM NaHS: 0.82  $\pm$ 310 0.08; 100µM NaHS: 0.84  $\pm$  0.07). These findings demonstrated increased H<sub>2</sub>S abundance 311 reduces hypoglossal neuronal activity consistent with findings using either ChrM459 or HO-2 312 null mice. Thus, the stability of inspiratory transmission from the preBötC to hypoglossal nucleus 313 appears to be negatively affected either by increasing H<sub>2</sub>S abundance or by disrupting HO-2 314 function.

315 CO produced by HO-2 is known to inhibit CSE-dependent  $H_2S$  production by HO-2 <sup>33,34</sup>. 316 Therefore, we sought to assess how CORM-3 (20µM), a pharmacological CO donor, impacted 317 activity in ChrMP459-treated wild type rhythmic slices (n= 3, Fig 6A) and rhythmic slices from 318 HO-2 null mice (n=4). Dysregulated HO-2 activity, caused by either pharmacological (ChrMP459) or genetic (HO-2 null mice) manipulation, is improved by CORM-3 as indicated by 319 320 larger I/O ratios (Fig 6B, dysregulated HO-2: 0.71 ± 0.04 vs. CORM-3: 1.03 ± 0.08, P=0.009) 321 and improved transmission (Fig 6C, dysregulated HO-2: 79.88 ± 4.69% vs. CORM-3: 322 93.39 ± 2.93%, P=0.036). As these findings suggested the absence of HO-2 dependent CO 323 production is a key factor driving transmission failure in the rhythmic slice preparation, we next 324 determined the involvement of CSE.

Inspiratory activity in the brainstem slice from HO-2:CSE double null mice appeared to be stable and consistent (**Fig 6D**). Quantification of simultaneous extracellular field recordings of preBötC activity and hypoglossal nucleus revealed a larger I/O ratio (**Fig 6E**, HO-2:CSE:  $1.04 \pm 0.02$ , n=6; P=0.003) and near absence of transmission failures (**Fig 6F**, HO-2:CSE:  $91.6 \pm 0.02\%$ ; P=0.0007) when compared to activity in HO-2 null slices.

Given the observations using HO-2:CSE null mice, we next sought to determine whether acute blockade of CSE could restore transmission relationships between the preBötC and the hypoglossal nucleus in the HO-2 null slice. *In vivo* L-PAG treatment improved transmission of preBötC activity to the hypoglossal nucleus in the rhythmic slice (**Fig. 7A,** n=6) as indicated by

334 larger cycle-to-cycle I/O ratios (Fig. 7B, L-PAG =  $1.01 \pm 0.03$ , P=0.008) and greater transmission 335 rates (Fig. 7C, L-PAG 96.31  $\pm$  2.62%, P<0.0001) when compared to the respective metrics from 336 untreated HO-2 null mice. Intermittent transmission failure was also evident in patch clamp 337 recordings from untreated HO-2 null hypoglossal neurons (Fig 7D, left shaded cycles) but not in 338 HO-2 null hypoglossal neurons treated with L-PAG (Fig. 7D, right). These reduced transmission 339 events correlated with smaller individual inspiratory drive currents in HO-2 null hypoglossal 340 neurons when compared to corresponding inspiratory drive currents from L-PAG treated HO-2 341 mice (Fig. 7D-E, HO-2: -36.71

 $\pm 2.14$  pA vs. L-PAG -194.3  $\pm 82.73$  pA, P=0.0007). Together, these experiments implicated the involvement of CSE-dependent H2S signaling with the effects of disrupted HO-2 signaling in hypoglossal neurons.

345 Blockade of small conductance calcium-activated potassium channel (SK<sub>Ca</sub>) activity 346 restores changes induced by HO-dysregulation in hypoglossal activity. As our 347 experiments implicated the involvement of H<sub>2</sub>S signaling, we next sought to determine how H<sub>2</sub>S 348 sensitive ion channels may contribute to impairing hypoglossal neuron activity caused by 349 HO-dysregulation.  $H_2S$  has been shown to enhance activity of several different potassium channels, including SK<sub>Ca</sub> and ATP-sensitive potassium channel (K<sub>ATP</sub>) activities <sup>35</sup>. As both SK<sub>Ca</sub> 350 and K<sub>ATP</sub> are important in the regulation of hypoglossal neuron excitability <sup>36,37</sup>, we examined 351 352 how blocking these channels affected hypoglossal activity during HO-dysregulation. At the network level, administration of the selective SK<sub>ca</sub> inhibitor, apamin (200µM), increased the 353 354 excitability of the hypoglossal neurons treated with ChrMP459. This enhanced activity exceeded 355 the original baseline activity (i.e., prior to ChrMP459 administration) causing ectopic bursting in 356 the hypoglossal nucleus (Supplemental Figure 2A) making analysis of population transmission 357 and I/O ratios unreliable. Therefore, we proceeded to resolve the effects of apamin on the 358 influence of ChrMP459 at the level of individual hypoglossal inspiratory motor neurons. While in 359 some hypoglossal neurons exposed to ChrMP459 apamin substantially increased drive currents

360 (>100pA; FIG 8A1), in others, apamin modestly increased the drive current (<100pA; FIG 8A2). 361 Despite this variability, apamin increased inspiratory drive currents received by ChrMP459 362 treated hypoglossal neurons (FIG 8A3, n= 6, ChrMP459: -85.77±38.54 pA vs. apamin: -363 219.97±97.76 pA, P=0.031). Apamin also enhanced the number of action potential generated 364 per preBötC burst during ChrMP459 (FIG 8B, n=8, ChrMP-459: 12.57 ± 3.7 action potential per 365 burst vs. apamin  $26.05 \pm 6.87$  action potential per burst, P=0.016). This was consistent with the 366 ability of apamin to reduce rheobase in ChrMP459 treated inspiratory hypoglossal neurons (FIG 367 8B; n=7, ChrMP459: 532.0 ± 186.5 pA vs. Apamin: 307.09 ± 80.9 pA, P=0.016).

368 To determine how blockade of  $K_{ATP}$  impacted hypoglossal activity during ChrMP459, we used 369 the  $K_{ATP}$  channel blocker, tolbutamide (100 $\mu$ M). Tolbutamide did not induce ectopic bursting in 370 the hypoglossal nucleus during ChrMP459 (Supplemental Fig 2B, n=5). Furthermore, 371 tolbutamide (100µM), did not improve the rate of transmission of preBötC activity 372 (Supplemental Fig 2C, ChrMP459: 69.24±6.0% tolbutamide: 71.23 ± 6.8%, P=0.76) but did 373 increase the I/O ratio (Supplemental Fig 2C right, ChrMP459: 0.68 ± 0.05 tolbutamide: 374 0.87 ± 0.05, P=0.037). Tolbutamide neither enhanced inspiratory drive currents in ChrMP459 375 (Supplemental Fig 2D, n=4, ChrMP459: -70.51 ± 27.49 pA vs. tolbutamide: -83.06 ± 36.29 pA, 376 P=0.37) nor increased the number of action potentials per preBötC burst in ChrMP459 (n=4, 377 ChrMP459: 7.063±2.08 action potential per burst vs. tolbutamide: 9.37±3.18 action potential 378 per burst, P=0.125). Moreover, tolbutamide did not affect rheobase of inspiratory hypoglossal neurons treated with ChrMP459 (Supplemental Fig 2D, n=4; ChrMP459: 221.01 ± 74.8 pA vs. 379 380 tolbutamide:  $180.4 \pm 68.63$  pA, P=0.218). Thus, these results suggested that apamin could 381 enhance activity of hypoglossal neurons during HO-inhibition; whereas, the efficacy of 382 tolbutamide to impact activity during HO-inhibition was limited.

### 383 Discussion

384 Our study reveals a previously uncharacterized neuromodulatory interaction between HO-2 and 385 CSE-derived H<sub>2</sub>S regulating activity from the hypoglossal nucleus. Our electrophysiological 386 studies revealed that dysregulated HO-2 activity impacted intrinsic properties of hypoglossal 387 neurons receiving drive from the preBötC. HO-2 dysregulation impaired transmission originating 388 from the rhythm generating preBötC and the motor nucleus, as evidenced by the cycle-to-cycle 389 reductions in the input-output relationship and the intermittent failures for preBötC activity to 390 generate corresponding motor pool output, which mitigated blocking CSE activity and mimicked 391 using a  $H_2S$  donor. The increased propensity of transmission failure in response to 392 dysregulated HO-2 activity and involvement of CSE / H<sub>2</sub>S are reminiscent of the OSA phenotypes observed in HO-2 null mice<sup>16</sup>. Thus, our study demonstrates the potential 393 394 importance of centrally derived interactions between HO-2 and CSE activity in regulating motor 395 neuron output important for maintaining upper airway patency.

396 Both a pharmacological inhibitor of HO and genetic elimination of Hmox-2 impaired transmission 397 of neural drive from preBötC to the hypoglossal nucleus. Mice treated with intermittent hypoxia 398 (IH) patterned after blood O<sub>2</sub> profiles seen during OSA also show failed transmission related to a changed input-output relationship between preBötC and hypoglossal nucleus<sup>31,38</sup>. While IH-399 400 induced phenomena are associated with alterations in preBötC activity, the HO inhibitor neither 401 impacted inspiratory rhythmogenesis from the preBötC (Fig. 1C) nor perturbed transmission of 402 neuronal drive to intermediate premotor neurons suggesting that changed interneuron 403 neurophysiology was not a major contributing factor to the impaired transmission in the 404 hypoglossal nucleus.

At the level of the hypoglossal nucleus, we documented that HO-2 is expressed among ChATpositive cells of the hypoglossal nucleus. Although we did not resolve expression among hypoglossal motor neurons innervating upper airway muscles, such as genioglossal neurons, our electrophysiological studies demonstrated that ChrMP459 had divergent effects on non-

inspiratory and inspiratory hypoglossal neurons. HO inhibition increased rheobase among noninspiratory neurons; whereas, in inspiratory hypoglossal neurons, it decreased the magnitude of drive currents, increased rheobase, and diminished the number of action potentials generated during preBötC bursting. Thus, while our studies did not resolve how hypoglossal neurons projecting to different tongue muscles are impacted by ChrMP459, our findings indicate that HO-dysregulation can differentially impact hypoglossal neurons that receive drive from the preBötC.

416 The observed postsynaptic phenomena among hypoglossal motor neurons receiving inspiratory 417 drive could contribute to the occurrence of obstructive apneas by increasing the probability for 418 intermittent reductions in upper airway tone from inspiratory drive by hypoglossal motor 419 neurons. Such loss of upper airway muscle tone can obstruct airflow even when inspiratory 420 activity is successfully produced at in other inspiratory muscles, such as the diaphragm. Indeed, this phenotype has been documented in HO-2 null mice<sup>16</sup>. However, further resolution is 421 422 needed to understand how dysregulation of HO-2 may impact hypoglossal neurons innervating 423 different muscle groups and to determine extent of the contribution from central HO-2 424 dysregulation to loss of upper airway tone and airway collapse in vivo.

In HO-2 null mouse, the incidence of OSA is absent with co-inhibition of CSE<sup>17</sup>, which is 425 consistent with reports that CO generated by HO-2 inhibits CSE-dependent H<sub>2</sub>S production<sup>33,34</sup>. 426 427 After documenting CSE expression in hypoglossal neurons and demonstrating an increased 428 abundance of H<sub>2</sub>S in the hypoglossal nucleus of HO-2 null mice, we demonstrated that a CO 429 donor improves transmission and the input-output relationship between the preBötC and the 430 hypoglossal nucleus in HO-2 null slices. Furthermore, using a  $H_2S$  donor also increased 431 transmission failures and reduced the I/O ratio similar to perturbing HO-2 activity. Endogenous 432  $H_2S$  activity could originate from other  $H_2S$  producing enzymes, such as cystathionone  $\beta$ synthase (CBS) that is expressed primarily in astrocytes throughout the CNS<sup>39</sup>. However, CBS 433 inhibition appears to have limited impact on inspiratory activity from the hypoglossal nucleus<sup>40</sup>. 434

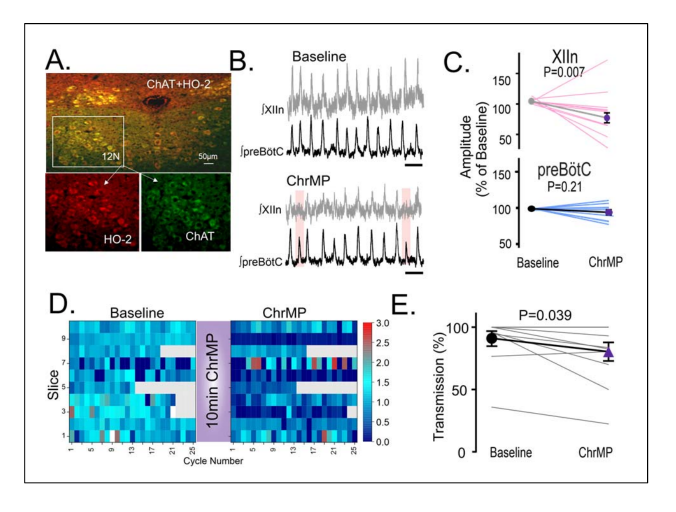
In contrast, our genetic and pharmacological manipulations to ablate/diminish CSE activity improved transmission and the input-output relationship between inspiratory drive and the hypoglossal nucleus in HO-2 null mice. Together these findings implicated the interaction between HO-2 / CO and CSE /  $H_2S$  within the hypoglossal nucleus to regulate its output.

439 How might enhanced H<sub>2</sub>S signaling reduce drive currents, and excitability of hypoglossal 440 neurons? While it is possible that H<sub>2</sub>S may impact presynaptic release of glutamate and/or 441 postsynaptic receptor activity of hypoglossal neurons, the ChrMP459 mediated increase in 442 rheobase among inspiratory hypoglossal neurons implicated the involvement of a non-synaptic 443 conductance(s) downstream of H<sub>2</sub>S based signaling caused by perturbations in HO-2 activity.  $H_2S$  can enhance both  $K_{ATP}$  and  $SK_{Ca}$  activities<sup>35</sup>. In the hypoglossal neurons,  $K_{ATP}$  is 444 dynamically active causing periodic adjustment of neuronal excitability<sup>36</sup> while SK<sub>Ca</sub> also 445 446 regulates excitability and firing properties of hypoglossal neurons<sup>37</sup>. In ChrMP459, tolbutamide 447 had a limited effect normalizing transmission as it improved the I/O ratio between preBötC and 448 the hypoglossal motor nucleus, but did not reduce transmission failure. At the neuronal level, 449 tolbutamide did not increase drive currents nor did it enhance intrinsic excitability of hypoglossal 450 neurons in ChrMP459. These results indicated that blockade of K<sub>ATP</sub> during HO-2 dysregulation 451 was limited in countering the effects on HO-2 dysregulation in the hypoglossal nucleus. On the 452 other hand, apamin normalized drive currents and increased excitability of inspiratory hypoglossal neurons in ChrMP459 demonstrating that blockade of SK<sub>Ca</sub> sufficiently mitigates 453 454 many aspects of HO-2 dysregulation in hypoglossal neurons that receive drive from the 455 preBötC.

In conclusion, our study provides proof-of-concept for the existence of a central mechanism by which loss of HO-2 leads to reduced upper airway tone by enhancing H<sub>2</sub>S activity in the hypoglossal nucleus. This mechanism appears to involve antagonistic interactions between HO-2 and CSE activities to regulate excitability of hypoglossal neurons and is localized in the neurons themselves. Although OSA in HO-2 null mice has attributed to increased "loop-

- 461 gain" arising from the hypersensitive carotid body reflex<sup>16</sup>, our findings indicate the potential
- 462 involvement of a disrupted interaction between HO-2 / CO and CSE / H<sub>2</sub>S in hypoglossal motor
- 463 neurons that contribute could to the sporadic loss of upper airway tone observed in OSA.

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464

465 Figure 1. Disruption of hemeoxygenase-2 (HO-2) in hypoglossal motor neurons impairs 466 inspiratory activity from the hypoglossal nucleus but not in the preBötC. A. HO-2 (red 467 bottom left) is expressed in ChAT<sup>+</sup> cells (green bottom right) of the hypoglossal nucleus (XIIn, 468 combined top). B-E. Population recordings of rhythmic brain slices were recorded from 469 ipsilateral preBotC and XIIn simultaneously, analyses were performed during baseline and 470 during bath application of 20µM ChrMP459 (ChrMP). B. Integrated traces of network activity in 471 spontaneously rhythmic brainstem slices (n=11) recorded from XIIn (grey) and preBötC (black) 472 before (top) and after (bottom) bath application, failed transmission events are highlighted (pink 473 boxes); scale bar: 5 sec. C. Percentage change of integrated burst amplitude from Baseline to 474 ChrMP in XIIn (top) and the preBötC (bottom). Thin pink and blue lines illustrate integrated burst amplitude from individual slices. Symbols connected by a thick line illustrate mean integrated
burst amplitude. **D.** Heat maps of consecutive cycle-to-cycle I/O ratios in Baseline and in
ChrMP. Each row represents a single slice experiment during baseline and in ChrMP. Grey
boxes indicate non-events in recordings from slower rhythms where less than 25 events (i.e., 25
cycles) occurred during the analysis window. **E.** Comparison of inspiratory drive transmission in
XIIn between Baseline and ChrMP. Thin grey lines illustrate transmission from individual slices.
Symbols connected by a thick line illustrate mean transmission.

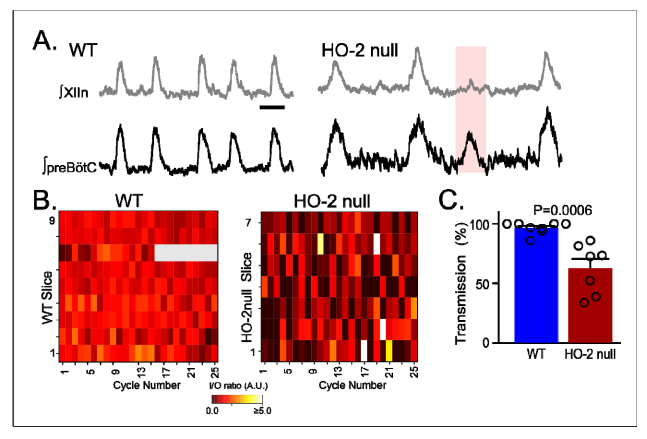
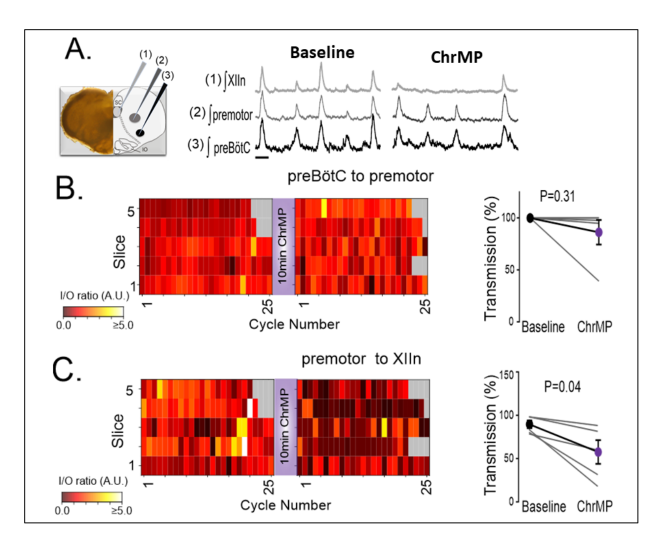


Figure 2. Genetic deletion of HO-2 reduces the I/O relationship between preBötC and the
 hypoglossal nucleus and uncouples of motor output from inspiratory rhythmogenesis.

A. Representative integrated traces of network burst activity in the preBötC and XIIn in rhythmic
slices from wildtype (WT) (left, n=9) and HO-2 null (HO-2<sup>-/-</sup>; right, n=7) mice; scale bar: 2 sec. B.
Heat maps of cycle-to-cycle I/O ratios in WT (left) and HO-2<sup>-/-</sup> (right) slices. Grey boxes indicate
non-events in recordings from slower rhythms where less than 25 events occurred during the
analysis window. C. Transmission of inspiratory activity from preBötC to XIIn in slices from WT
vs HO-2<sup>-/-</sup>. Symbols illustrate transmission from individual slices.

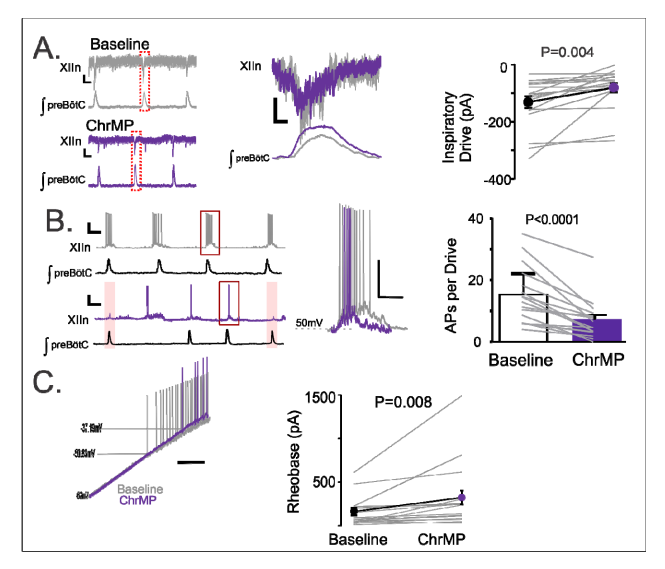
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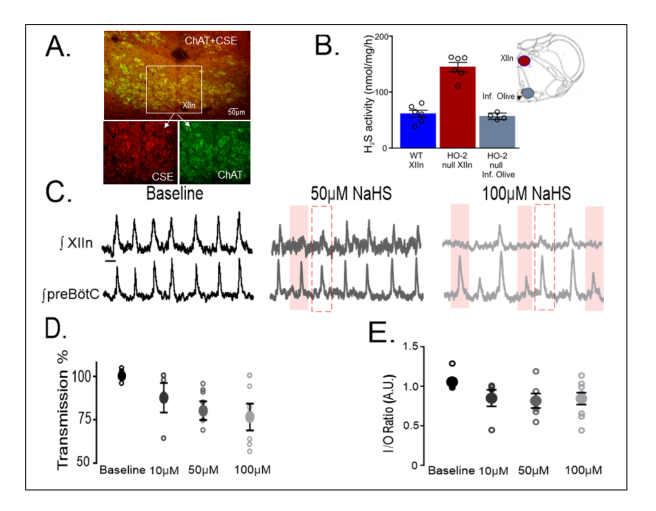
Figure 3. While ChrMP459 does not change transmission from the preBötC to the 493 494 premotor area, ChrMP459 increases transmission failure from the premotor area to the 495 hypoglossal nucleus. A. Diagram of medullary brain slice illustrating relative electrode 496 placement for simultaneous triple extracellular recordings (n=5) from the XIIn (light grey, 1), 497 premotor field (dark grey, 2), and preBötC (black, 3). Corresponding representative traces of 498 integrated network activity in Baseline (left) and in 20µM ChrMP (right); scale bar: 1 sec. B. 499 Heat maps of the cycle to cycle I/O ratio (left) and transmission (right) between preBötC and the 500 premotor field. C. Heat maps of the cycle to cycle I/O ratio (left) and transmission (right) 501 between the premotor field and XIIn.

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502 Figure 4. Heme oxygenase inhibition reduces inspiratory drive currents in hypoglossal 503 neurons. Whole cell patch clamp recordings were made from hypoglossal neurons in rhythmic 504 brain slices while simultaneously recording ipsilateral preBötC activity in Baseline and in ChrMP. 505 Neurons were disinhibited from fast synaptic inhibition using 50µM PTX and 1µM Strychnine. 506 A. (left) Representative voltage clamp recordings from a XIIn neuron (V<sub>holding</sub>= -60mV) aligned 507 with corresponding integrated network activity from preBötC before (Baseline, top, grey) and 508 after 20 µM ChrMP (bottom, purple). Scale bar: 1 sec x 10pA. (middle) Magnification of 509 highlighted (red dotted box) drive currents from Baseline (grey) and ChrMP (purple). Scale bar: 510 100 msec x 10pA. (right) Comparison of XIIn inspiratory drive current magnitude in Baseline and

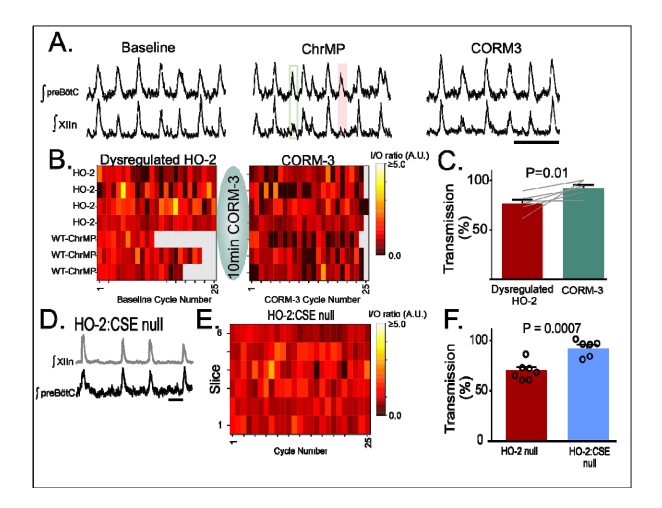
511 ChrMP (n=19). Thin grey lines illustrate individual neuron response. Symbols connected by a 512 thick line illustrate mean drive current. B. (left) Representative current clamp recordings from a 513 XIIn neuron spontaneously active with the preBötC network rhythm in Baseline (top, grey) and 514 20 µM ChrMP (bottom, purple); skipped transmission of action potentials in ChrMP are 515 highlighted (pink). Scale bar 2sec x 20mV. (middle) Magnification of highlighted neuronal activity 516 (red box in left). Scale bars: 100msec x 25mV. (right) Number of action potentials generated 517 per inspiratory burst in Baseline and in ChrMP (n=17). Thin grey lines illustrate individual neuron 518 response. C. (left) Representative trace of current clamp recording in response to ramp current 519 injection during Baseline (grey trace) and in ChrMP (purple trace); scale bar: 500 msec. (right) 520 Comparison of rheobase in inspiratory XIIn neurons during Baseline and in ChrMP (n=19). Thin 521 grey lines illustrate individual neuron response.



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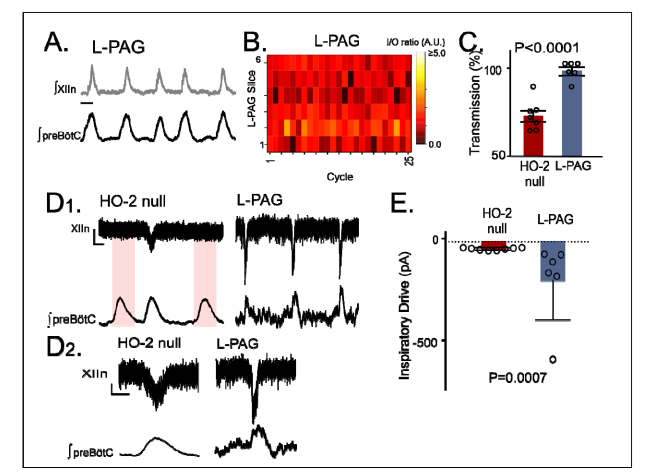
524 Figure 5. CSE-dependent  $H_2S$  is produced in the hypoglossal nucleus and exogenous 525 NaHS uncouples hypoglossal nucleus activity from the preBötC. A. CSE (red, bottom left) 526 is expressed in ChAT<sup>+</sup> neurons (green, bottom right) in the XIIn (overlay, top). B. CSEdependent  $H_2S$  generation in homogenates from WT and HO-2<sup>-/-</sup>. Homogenates were prepared 527 528 from tissue punches from the XIIn (red area in slice diagram) and inferior olive (grey area in slice diagram) at bregma between -7.20mm and -7.90mm. (WT: XIIn n=6; HO-2<sup>-/-</sup>: XIIn n=6, 529 530 inferior olive n=4). Note: Each n value reported in B represents a sample consisting of the anatomical region pooled from two animals. C. Integrated traces from XIIn (top) and preBötC 531 532 (bottom) during baseline (black), and in response to the  $H_2S$  donor, NaHS, at 50  $\mu$ M (dark grey) 533 and 100 µM (light grey). After NaHS application, XIIn but not preBötC burst amplitude were 534 diminished (red dashed box) and in some cases, preBötC drive failed to produce activity in the bioRxiv preprint doi: https://doi.org/10.1101/2022.03.23.485481; this version posted August 10, 2022. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

- 535 XIIn (pink boxes). **D.** Comparison of transmission from preBötC to XIIn after NaHS application at
- 536 10µM, 50µM and 100µM. E. I/O ratios for each NaHS concentration. (Baseline: n=9; 10µM n=5;
- 537 50µM n=6; 100µM n=9).



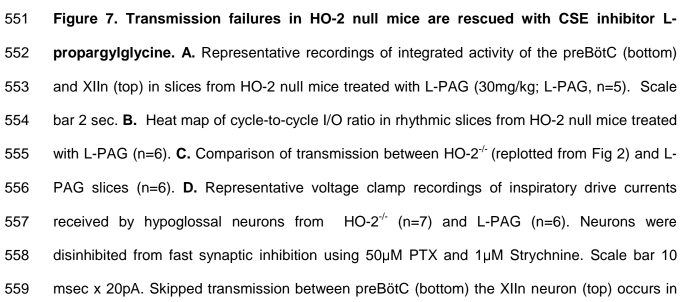
538 Figure 6. HO-dependent transmission failures can be recovered with CO-donor CORM-3 and are not present in HO-2:CSE null transmission. A. Representative extracellular 539 540 recording from preBotC and XIIn in WT slices during Baseline (left), in ChrMP alone (middle), 541 and in ChrMP + 20uM CORM-3 (CORM3, right). B. Heat maps of cycle-to-cycle I/O ratios 542 during dysregulated HO-2 (n=7: n=4 HO-2 null and n=3 WT-ChrMP) before and after CORM-3 543 application. C. Transmission from preBötC to XIIn from dysregulated HO-2 slices before and after bath application of CORM-3. D. Representative extracellular recordings from preBötC and 544 XIIn in slices from HO-2:CSE null (HO-2:CSE<sup>-/-</sup>); scale bar 2 sec. **E.** Heat map of cycle-to-cycle 545 I/O ratio from preBötC to XIIn in HO-2:CSE<sup>-/-</sup>. The I/O ratio from HO-2:CSE<sup>-/-</sup> is greater than I/O 546 ratios from HO-2<sup>-/-</sup> (P=0.003). F. Comparison of transmission from preBötC to XIIn in HO-2<sup>-/-</sup> 547

## and HO-2:CSE<sup>-/-</sup> (n=6). HO- $2^{-/-}$ data used for comparisons in E and F were originally shown in



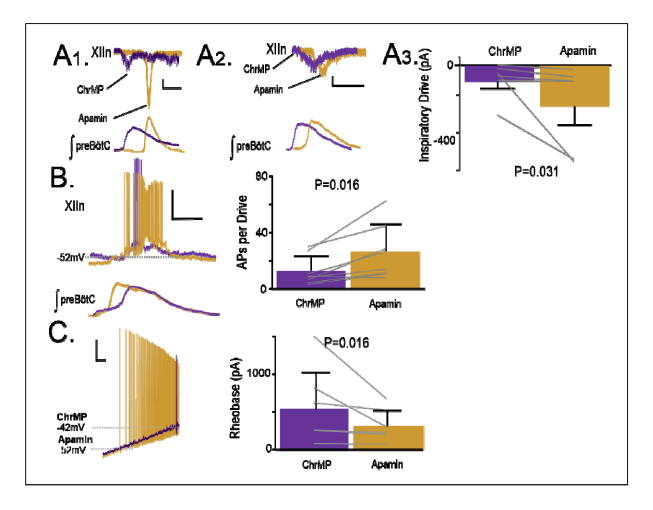
549 Figure 2.





560 untreated HO-2<sup>-/-</sup> (highlighted by pink boxes) but not in neurons from L-PAG. **D1.** 561 Magnified representative drive potentials from HO-2<sup>-/-</sup> and L-PAG (from highlighted by red-562 outlined boxes in **D**). scale bars: 100 msec x 10pA. **E.** Quantification of drive currents received 563 by XIIn motoneurons from HO-2<sup>-/-</sup> mice (n=8) produce smaller drive potentials when compared 564 to L-PAG (n=6).

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566

567 Figure 8. Apamin improves network input-output relationship between hypoglossal 568 nucleus and reverses changes to intrinsic and synaptic excitability caused by HO 569 **dysregulation.** A1. Representative inspiratory drive current received by hypoglossal neurons in 570 ChrMP (purple) and in Apamin (200  $\mu$ M, gold) where apamin increased the drive current >100 571 pA. Scale bars: 1 s x 50 pA. A2. Representative inspiratory drive current hypoglossal neuron in 572 ChrMP (purple) and in Apamin (gold) where apamin increased the drive current by less than 573 <100 pA Scale bars: 1 s x 50 pA. A3. Comparison of inspiratory drive currents from 574 hypoglossal neurons exposed to ChrMP (200 µM) then treated with Apamin (n=6). The effect 575 of ChrMP on baseline disinhibited drive current for each of these neurons were reported in Fig 576 4A. B. (left) Representative trace of bursting of a hypoglossal neuron resulting from preBötC 577 drive during ChrMP (purple) and in Apamin (gold) (n=8). Scale bars: 20 mV x 500 msec. (right)

578 Comparison of action potentials generated per preBötC burst during ChrMP and Apamin (n=8).

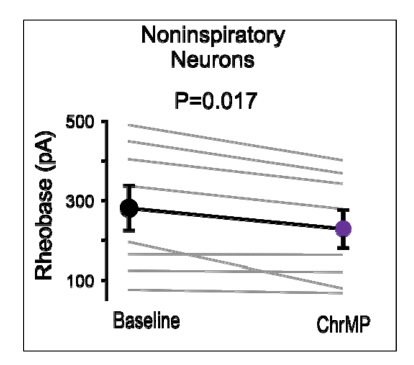
579 The effect of ChrMP on baseline action potential generated per preBötC burst for each of these

580 neurons were reported in Fig 4B. C. (left) Representative traces of current clamp recordings in

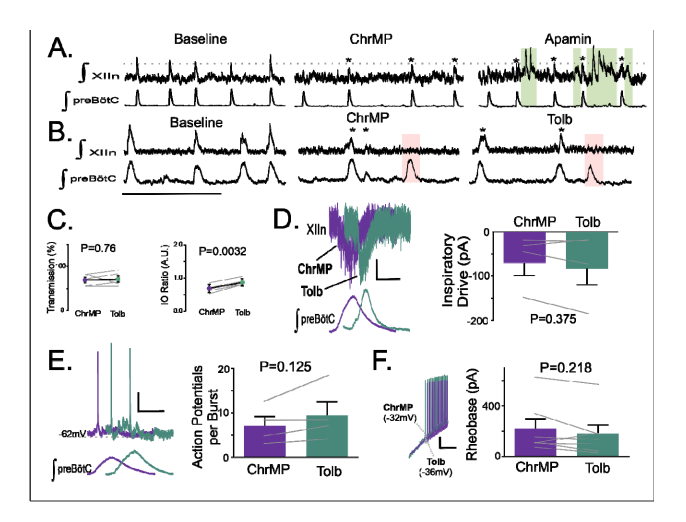
response to ramp current injection in ChrMP (purple) and in Apamin (gold). Scale bar: 500msec

- 582 x 10mV). (right) Rheobase comparison from inspiratory XIIn neurons during ChrMP and Apamin
- 583 (n=7). The effect of ChrMP on baseline rheobase for each of these neurons were reported in
- 584 Fig 4C.
- 585

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Supplemental Figure 1: ChrMP459 decreases rheobase of non-inspiratory hypoglossal neurons. Comparison of rheobase from non-inspiratory hypoglossal neurons in baseline and ChrMP (n=8). Non-inspiratory hypoglossal neurons were defined as neurons that did not receive synaptic drive in-phase with a preBötC burst. Neurons were disinhibited from fast synaptic inhibition using 50µM PTX and 1µM Strychnine.



591 Supplemental Figure 2: Apamin produce ectopic hypoglossal network activity during 592 ChrMP459 and tolbutamide has limited effects on network and inspiratory hypoglossal 593 neurons activity during ChrMP459. A. Representative traces of integrated network activity of 594 the preBötC (top) and XIIn (bottom) in slices from WT mice Baseline, ChrMP, and in 200 µM 595 Apamin. Asterisks (\*) indicate detected hypoglossal output in phase with the preBötC. Green 596 box indicates ectopic network bursting. Scale bar 10 sec. Due to ectopic bursting within and 597 around rhythmic burst transmissions, analysis of apamin at the network level could not be 598 accurately detected. **B.** Representative traces of integrated network activity of the preBötC (top) 599 and XIIn (bottom) in slices from WT mice Baseline, ChrMP, and in tolbutamide (200  $\mu$ M, Tolb).

600 Asterisks (\*) indicate detected hypoglossal output in phase with the preBötC. Pink box indicates 601 cycles where preBötC drive failed to produce activity in the XIIn. C. Comparisons of 602 Transmission (left) and I/O ratio (right) in ChrMP459 and in Tolbutamide (n=5). D. (top) 603 Representative traces of inspiratory drive currents in ChrMP (purple) and in Tolb (green). Scale 604 bar 1 sec x 50 pA. (bottom) Comparison disinhibited inspiratory drive currents in ChrMP and 605 Tolb (n=4). The effect of ChrMP on baseline drive current for each of these neurons were 606 reported in Fig 4B. E. (left) Representative trace of bursting of a hypoglossal neuron resulting 607 from preBötC drive during ChrMP (purple) and in Tolb (green) (n=4). Scale bars: 10 mV x 608 500 msec. (right) Comparison of action potentials generated per preBötC burst during ChrMP 609 and Tolb (n=4). The effect of ChrMP on baseline action potentials generated per preBötC burst 610 for each of these neurons were reported in Fig 4C. F. (left) Representative traces of current 611 clamp recordings in response to ramp current injection in ChrMP (purple) and in Tolb (green). 612 Scale bar: 1 sec x 10mV). (right) Rheobase comparison from inspiratory hypoglossal neurons 613 during ChrMP and Tolb (n=7). The effect of ChrMP on rheobase for each of these neurons were 614 reported in Fig 4C.

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