

1 **Title:** Type-I nNOS neurons orchestrate cortical neural activity and vasomotion
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21 **Abstract:** It is unknown how the brain orchestrates coordination of global neural and vascular dynamics. We sought
22 to uncover the role of a sparse but unusual population of genetically-distinct interneurons known as type-I nNOS
23 neurons, using a novel pharmacological strategic to unilaterally ablate these neurons from the somatosensory cortex
24 of mice. Region-specific ablation produced changes in both neural activity and vascular dynamics, decreased power
25 in the delta-band of the local field potential, reduced sustained vascular responses to prolonged sensory stimulation,
26 and abolished the post-stimulus undershoot in cerebral blood volume. Coherence between the left and right
27 somatosensory cortex gamma-band power envelope and blood volume at ultra-low frequencies was decreased,
28 suggesting type-1 nNOS neurons integrate long-range coordination of brain signals. Lastly, we observed decreases
29 in the amplitude of resting-state blood volume oscillations and decreased vasomotion following the ablation of type-
30 I nNOS neurons. This demonstrates that a small population of nNOS-positive neurons are indispensable for
31 regulating both neural and vascular dynamics in the whole brain and implicates disruption of these neurons in
32 diseases ranging from neurodegeneration to sleep disturbances.
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35 **Keywords:** neurovascular coupling, type-I nNOS neurons, somatosensory cortex, two-photon, cerebral blood
36 volume, widefield optical imaging, sleep

37 **Introduction**

38 Coordinated neural and hemodynamic activity across the brain are vital for arousal and cognition. The
39 relationship between changes in neural activity and (typically measured) changes in blood volume, flow, and
40 oxygenation, which are frequently studied with hemodynamic imaging and form the foundation of modern
41 neuroscience, are mediated by local vasodilation and is known as neurovascular coupling (Drew, 2019a; Schaeffer,
42 2021). The vasodilator nitric oxide (NO) (Hosford, 2019), produced by neuronal nitric oxide synthase (nNOS) in
43 neurons, provides a key mechanism of arteriole dilation. Neurons in somatosensory cortex that express nNOS are
44 activated by sensory stimulation, volitional whisking, and locomotion (Echagarruga, 2020; Ahn, 2023), all of which
45 produce robust hemodynamic responses (Huo, 2014; Winder, 2017). Chemogenetic elevation or suppression of
46 nNOS neuron activity alters both baseline diameter and behaviorally-evoked dilation of arterioles (Echagarruga,
47 2020). Similar vascular changes are produced by local infusion of a NOS inhibitor (Echagarruga, 2020), implicating
48 NO signaling directly in vascular regulation. Optogenetic activation of nNOS-positive neurons produces varying
49 effects, with some reports demonstrating vasodilation and blood flow increases with minimal changes in neural
50 activity (Krawchuk, 2020; Lee, 2020; Ruff, 2024), while others show large, low-frequency EEG changes during sleep
51 and quiet wakeful states (Gerashchenko, 2018). While NO is known to modulate the excitability of neurons (Kara,
52 1999; Smith, 2003), artificial activation of nNOS neurons seems to drive vascular responses that are disproportionately
53 larger than any corresponding neural changes, suggesting that this small population of neurons has an impact on
54 vascular signals that is substantially greater than that of overall neural activity.

55 Two distinct GABAergic interneurons types express nNOS, denoted as type-I and type-II (Kawaguchi, 1997;
56 Kubota, 2011; Perrenoud, 2012). nNOS-positive neurons make up approximately 20% of GABAergic interneurons in
57 the cortex (Hendry, 1987; Sahara, 2012), with type-I constituting only 0.5–2% depending on brain region (Yan, 1997;
58 Tricoire, 2012; Chong, 2019). Type-I nNOS neurons are sparse, somatostatin-positive, most dense in the deep layers
59 of cortex, and express nNOS at higher levels than type-II. They extend both local and long-range projections
60 throughout the cortex, receive excitatory input from nearby pyramidal neurons and drive increases in cerebral blood
61 flow when stimulated (Ruff, 2024). Type-II nNOS neurons are in contrast much more abundant, smaller in size and
62 morphology, and express nNOS less strongly than type-I. Type-II are a more heterogeneous group that express a
63 variety of interneuron subtype markers and are more evenly distributed throughout the different layers of cortex
64 (Yan, 1996; Perrenoud, 2012). Type-I nNOS neurons have been observed in rodents, carnivores, and primates
65 (Tomioka, 2005; Higo, 2007; Tomioka, 2007) and are active during sleep (Gerashchenko, 2008; Kilduff, 2011;
66 Morairty, 2013; Dittrich, 2015). In the cortex, type-I nNOS neuron density is anticorrelated with vascular density
67 (Wu, 2022). Type-I nNOS neurons integrate local activity from feedforward excitatory pathways from the cortex and
68 thalamus, likely contributing to the correlated changes in blood flow that are observed between functionally
69 connected regions across hemispheres (Ruff, 2024). Molecular, electrophysiological, and immunohistochemistry
70 studies have demonstrated that type-I nNOS neurons are the only cells in the cortex that express the tachykinin
71 receptor 1 (also known as neurokinin 1 receptor, TACR1/NK1R), the primary receptor for the endogenous

72 neuropeptide substance P (Penny, 1986; Vruwink, 2001; Kubota, 2011; Dittrich, 2012; Endo, 2016; He, 2016; Huang,
73 2016; Paul, 2017; Vanlandewijck, 2018; Matsumura, 2021; Ruff, 2024). Local infusion of substance P into the cortex
74 causes a sustained increase in basal arterial diameter that is dependent upon local neural activity (Echagarruga,
75 2020). Type-I nNOS neurons likely receive substance P from the ~40% of parvalbumin (PV)-positive cortical
76 interneurons (Vruwink, 2001; Pfeffer, 2013; Bugeon, 2022) that are thought to drive network synchrony (Cardin,
77 2009; Sohal, 2009). Optogenetic stimulation of PV neurons for several seconds drives a biphasic hemodynamic
78 response, comprised of an early constriction (driven by suppression of overall population activity through GABA
79 release), and a delayed, prolonged dilation lasting tens of seconds that is mediated indirectly by substance P (Vo,
80 2023). The delayed vasodilation was not directly related to pyramidal neuron activity, was blocked by TACR1
81 antagonists, and was occluded by substance P, suggesting that activation of PV neurons drives downstream
82 activation of type-I nNOS neurons. These experiments point to type-I nNOS neurons as having a large effect on
83 vascular dynamics despite their smaller contribution in driving overall neural activity. However, optogenetic
84 stimulation only activates one component of a circuit in isolation, in an otherwise healthy and intact microcircuit,
85 potentially giving an incomplete picture of the neuron's role. Type-I nNOS neurons are uniquely vulnerable to stress
86 (Han, 2019), and their loss is likely in part causal to neurodegeneration (Schaeffer, 2021; Iadecola, 2023), making it
87 vital to understand brain wide changes following a more physiologically-relevant model of perturbation.

88 Here we sought to reveal the role of type-I nNOS neurons in controlling neural and vascular dynamics in the
89 cortex by selectively ablating them with saporin conjugated to a substance P analog (SP-saporin). Using this targeted
90 ablation approach, we found that ablation of type-I nNOS neurons caused decreases in the hemodynamic response
91 to sustained vibrissae stimulation, eliminated the post-stimulus undershoot, decreased local field potential power
92 in the delta-band (1-4 Hz), reduced bilateral correlations in gamma-band power and blood volume across arousal
93 states, and reduced the amplitude of resting-state blood volume oscillations. Together, these experiments
94 demonstrate that a small subset of type-I nNOS neurons regulate key neural and vascular dynamics that coordinate
95 changes in vasomotion.

96 **Results**

97 We investigated the effects of localized ablation of type-I nNOS neurons in somatosensory cortex on
98 neurovascular coupling and functional connectivity in unanesthetized, head-fixed mice. C57BL6J mice (119 total,
99 both male and female) were injected with either saporin conjugated to a substance P analog with high affinity for
100 the substance P receptor (SP-SAP) or a scrambled peptide as a control (Blank-SAP) into a localized region of one
101 hemisphere's somatosensory cortex. We used widefield optical imaging, electrophysiology, and 2-photon
102 microscopy to evaluate neural and hemodynamic changes following targeted ablation of type-I nNOS neurons in the
103 vibrissae (whisker) representation of somatosensory cortex while carefully monitoring arousal state (see **Materials**
104 and **Methods**). All imaging was performed during the animals' light cycle. Statistics included generalized linear

105 mixed-effects model (GLME), general linear models (GLM), or unpaired t-tests with corrections for multiple
106 comparisons when necessary.

107 **Saporin-conjugated peptides produce selective targeted ablation of type-I nNOS neurons**

108 While optogenetic and chemogenetic models give us insights into the potential function of neuron subtypes, it
109 is difficult to completely silence neurons *in vivo*, and the patterns of activity they induce are not physiological and
110 can have paradoxical effects on neural activity (Li, 2019; Andrei, 2021). Saporin provides a pharmacological route to

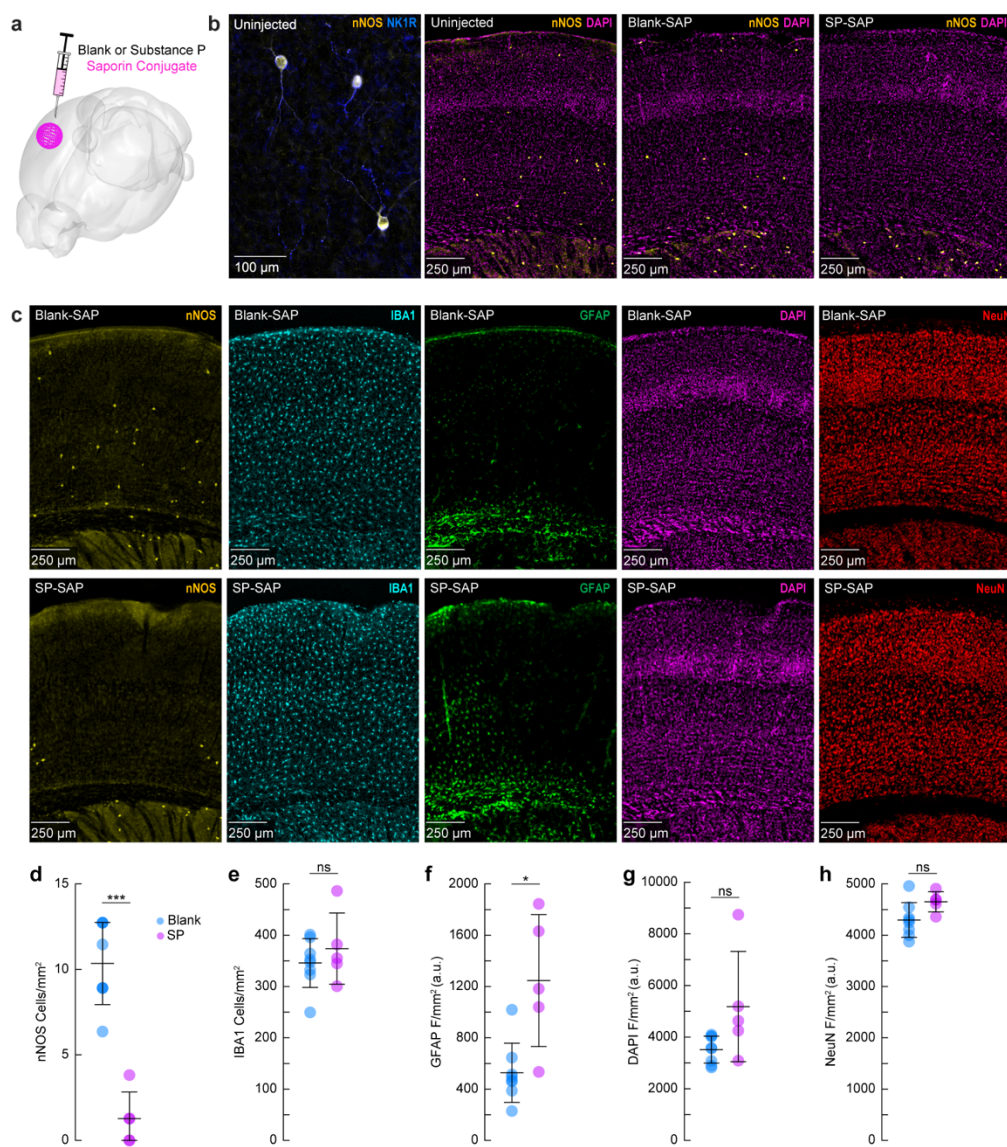


Figure 1. Targeted ablation of type-I nNOS neurons with cortical SP-SAP injections (a) Schematic showing intracortical administration of the ribosome inactivating protein saporin conjugated to either SP or a vehicle control. (b) Representative immunofluorescence of anti-nNOS (yellow) and anti-NK1 (blue) showing colocalization of the NK1 receptor on cortical nNOS-positive neurons. Representative immunofluorescence of anti-nNOS (yellow) and DAPI (magenta) from animals with no injection (Uninjected, left), Blank-SAP (middle), or SP-SAP (right). (c) Representative immunofluorescence of Blank-SAP (top) and SP-SAP (bottom) sections co-stained with nNOS (left), IBA1 (middle left), GFAP (middle), DAPI (middle right), or NeuN (right). NeuN image was taken from section immediately adjacent to the first four. (d) Quantification of nNOS counts, (e) IBA1 counts, (f) GFAP fluorescence, (g) DAPI fluorescence, (h) NeuN fluorescence. Error bars (d-h) denote SD. Bonferroni correction (5) * α < 0.01, ** α < 0.002, *** α < 0.002. k-l * α < 0.05, ** α < 0.01, *** α < 0.001, GLME.

111 selectively kill only the cells that internalize it, and saporin conjugated to a peptide effectively targets neurons that
112 express a receptor for the peptide (McKay, 2008; Abbott, 2012), allowing targeting of specific cell types orthogonal
113 to any genetic targeting techniques.

114 We first sought to validate the efficacy and specificity of saporin-based targeting of cortical TACR1-expressing
115 neurons. Type-I nNOS neurons were ablated by injection of the ribosome inactivating protein saporin (SAP),
116 conjugated either to substance P (SP-SAP) or a scrambled peptide as a control (Blank-SAP) (**Fig. 1a**). The SP-bound
117 SAP toxin binds to TACR1-expressing neurons (Wiley, 1999; Martin, 2001; Wang, 2002), which in the cortex are
118 exclusively expressed by type-I nNOS neurons (**Fig. 1b**). We used both immunofluorescence (**Fig. 1c**) and NADPH
119 diaphorase staining (a histochemical marker for NOS, **Fig. S1**), to visualize nNOS neuron ablation (Hope, 1989; Bredt,
120 1991; Hope, 1991). Immunofluorescent labeling in mice injected with Blank-SAP showed labeling of nNOS-positive
121 neurons near the injection site. In contrast, mice injected with SP-SAP showed a clear loss in nNOS-labeling around
122 the injection site, with nNOS-positive neurons both subcortically and in the entirety of the contralateral hemisphere
123 remaining intact. We quantified the efficacy of SP-SAP in removing type-I nNOS neurons by quantifying the number
124 of nNOS-positive neurons per square mm of cortical tissue (**Fig. 1d**), of which SP-SAP injected mice ($N = 5$, 3M/2F)
125 had a significant reduction of nNOS-labeled neurons (1.3 ± 1.6 neurons/mm 2) compared to Blank-SAP mice ($N = 8$,
126 4M/4F) with 10.3 ± 2.4 neurons/mm 2 (GLME $p = 5.74 \times 10^{-6}$).

127 We checked for non-specific effects driven by SP-SAP injections one-month post-administration by staining for
128 ionized calcium binding adaptor molecule 1 (IBA1), glial fibrillary acidic protein (GFAP), DAPI, and NeuN (neuronal
129 nuclei) as well as nNOS (as a positive control for ablation). There was no difference in the number of microglia (**Fig.**
130 **1e**) between groups (Blank-SAP: 345.8 ± 47.7 microglia/mm 2 ; SP-SAP: 373.8 ± 69.4 microglia/mm 2). There was an
131 increase in GFAP labeling following SP-SAP injection (1246.3 ± 514.8 AU/mm 2 ; Blank-SAP: 527.2 ± 230.9 AU/mm 2 ,
132 GLME $p = 0.0029$) (**Fig. 1f**). There was no significant difference in DAPI fluorescence (Blank-SAP: 3511.2 ± 524.42
133 AU/mm 2 ; SP-SAP: 5180.7 ± 2135.0 AU/mm 2 ; GLME $p = 0.038$, **Fig. 1g**) or NeuN fluorescence (Blank-SAP: $4296.8 \pm$
134 339.4 AU/mm 2 ; SP-SAP: 4652.1 ± 196.8 AU/mm 2 ; GLME $p = 0.043$, **Fig. 1h**) between groups after Bonferroni
135 correction for (5) multiple comparisons (**Fig. 1d-h**). Together, these findings indicate that the SP-SAP toxin was highly
136 selective in ablating type-I nNOS neurons with minimal non-specific effects. All mice that underwent imaging were
137 histologically verified for successful type-I nNOS neuron ablation using NADPH diaphorase staining. We quantified a
138 subset of these mice and saw a similar specific removal of type-I nNOS neurons (**Fig. S1**). To determine if ablation of
139 type-I nNOS neurons from the somatosensory cortex had any impact on behavior or arousal state, we assayed
140 exploratory behavior, sleep, and pupil dynamics (**Fig. S2**). To assay exploratory behavior, mice were placed in a novel
141 open field environment and allowed to explore for 5 minutes while quantifying distance traveled and time spent in
142 the center, two metrics for evaluating stress and anxiety in rodents (Roth, 1979; Seibenhener, 2015). We noted no
143 differences in any metric evaluating open field behavior ($N = 12-23$ mice/group). We also assessed multiple
144 measurements of sleep quality and quantity. There was no difference in the percentage of time each mouse spent

145 in rapid eye movement (REM) or non-REM (NREM) sleep, or the percentage of time they spent volitionally whisking
146 while awake (N = 9 mice/group). We also noted no significant differences in eye-related arousal state metrics
147 including interblink-interval, pupil size, and pupillary response to vibrissae stimulation (Turner, 2023). Together, this
148 suggests that ablation of type-I nNOS neurons had no gross effects on sleep or general ambulatory behavior.

149 **Impact of type-I nNOS neuron removal on neural and hemodynamic signals across arousal states**

150 To determine the impact localized ablation of type-I nNOS neurons had on neural and hemodynamic signals in
151 the somatosensory cortex, we used widefield optical imaging (Sirotin, 2009; Huo, 2014) to measure changes in total
152 hemoglobin ($\Delta[\text{HbT}]$, an indicator of blood volume). We measured neural activity using either implanted electrodes
153 (Winder, 2017) to measure changes in local field potential (LFP) (Fig. 2a, c, d), or in a separate cohort of mice, pan-
154 neuronal expression of the calcium indicator GCaMP7s (Chan, 2017; Dana, 2019), which provides complementary
155 measures of bulk neural activity (Fig. 2b, e, f). In mice expressing GCaMP7s, we measured $\Delta[\text{HbT}]$, as well as changes

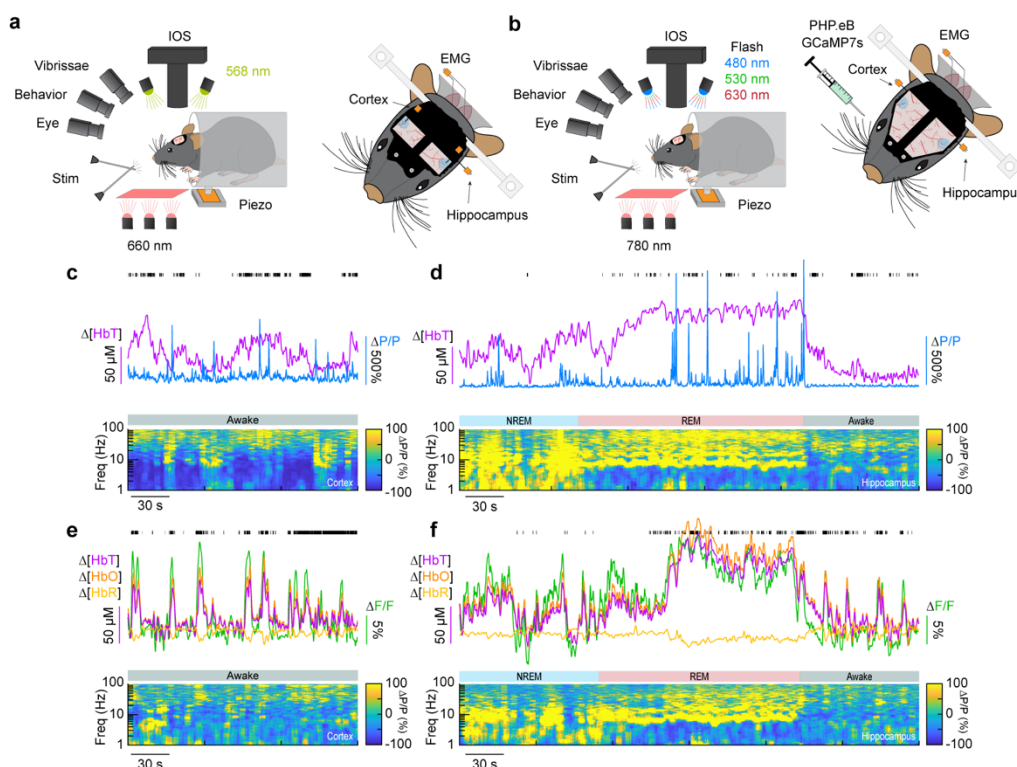


Figure 2. Simultaneous measurement of neural and hemodynamic signals in mice across arousal states (a) Schematic of widefield optical imaging experimental setup. The brain is illuminated at an isosbestic wavelength of hemoglobin (568 nm). Changes in reflected light measuring changes in total blood volume are captured by a camera mounted above the head while several other cameras monitor animal behavior and arousal state including vibrissae and pupil tracking. Vibrissae stimulation is done by directed air puffs. Polished and reinforced thinned-skull windows were bilaterally implanted over the somatosensory cortex. Tungsten stereotrodes were implanted underneath each window to record changes in local field potential within the area of interest. An additional hippocampal stereotrode and a neck electromyography electrode were used to assist in sleep scoring. (b) Schematic of widefield optical imaging experiments with GCaMP7s. Alternating illumination with 480 nm, 530 nm, and 630 nm light were used to measure changes in total hemoglobin, blood oxygenation, and GCaMP7s fluorescence. (c, e) During the awake state, power in low-frequency cortical LFP is low and power in the gamma-band (30-100 Hz) is elevated during activity such as volitional whisking. (d, f) Periods of NREM and REM sleep are accompanied by large oscillations in cerebral blood volume, with large increases in power in delta-band (1-4 Hz) cortical LFP during NREM and large increases in theta-band (4-10 Hz) hippocampal LFP during REM.

156 in cerebral oxygenation ($\Delta[\text{HbO}]$) and deoxygenation ($\Delta[\text{HbR}]$) using alternating illumination at 480/530/630 nm (Ma,
157 2016b; Ma, 2016a; Zhang, 2019). We corrected for hemodynamic contamination of GCaMP7s signals using the
158 simultaneously acquired hemoglobin signals (Kramer, 1979; Ma, 2016b; Wright, 2017; Scott, 2018). Measurements
159 of neural and hemodynamic signals were taken bilaterally through polished and reinforced thinned-skull windows
160 (Drew, 2010; Shih, 2012) in the vibrissae representation of somatosensory cortex. Each animal was habituated to
161 head-fixation over the course of several days following surgery. For the first 60 minutes of each recording session,
162 the vibrissae were briefly stimulated with directed puffs of air to either the left or right pad, or by a puffer directed
163 away from the mouse as an auditory control (Drew, 2011). Afterwards, each mouse was given several hours to
164 naturally sleep with no stimulation. To determine the arousal state of the mouse, we performed electromyography
165 of the nuchal muscles of the neck, tracked vibrissae movement and pupil diameter using video, and body movements
166 with a force sensor. Arousal state was scored in 5 second intervals as either Awake, NREM, or REM as previously
167 described (Turner, 2020) from behavioral and physiological data using a bootstrapped random forest classification
168 algorithm. We saw no differences in the accuracy or validity of our sleep-scoring models across different
169 experimental conditions (Blank-SAP; SP-SAP; Uninjected) (Fig. S3).

170 Ablation of type-I nNOS neurons reduces the stimulus-evoked response

171 We first determined the impact of removal of type-I nNOS neurons on evoked hemodynamic signals. The initial
172 increase in blood volume in response to brief (0.1 second) stimulation of the contralateral vibrissae was not affected
173 by ablation of type-I nNOS neurons (Fig. 3a), but the post-stimulus undershoot was absent in the SP-SAP mice. When
174 we evaluated the canonical post-stimulus hemodynamic undershoot from 2:4 seconds, the Blank-SAP group ($N = 9$,
175 5M/4F) had a mean of $-2.2 \pm 0.5 \mu\text{M}$, compared to the SP-SAP group ($N = 9$, 5M/4F) mean of $1.0 \pm 0.6 \mu\text{M}$ (GLME p
176 = 0.0005). This result is consistent with the observation that type-I nNOS neurons express the vasoconstrictory
177 neuropeptide Y (NPY) (Karagiannis, 2009) which is thought to underlie this post-stimulus undershoot (Uhlirova,
178 2016). It is also possible that loss of NO signaling from type-I nNOS neurons could also contribute through
179 interactions with blood volume to generate oscillations and post-dilation undershoots (Haselden, 2020). Using three-
180 wavelength spectroscopy in mice expressing GCaMP7s, we saw consistent increases in blood volume, neural activity,
181 and cerebral oxygenation during prolonged (5 seconds) vibrissae stimulation. $\Delta[\text{HbT}]$ evaluated 1.5:6.5 seconds

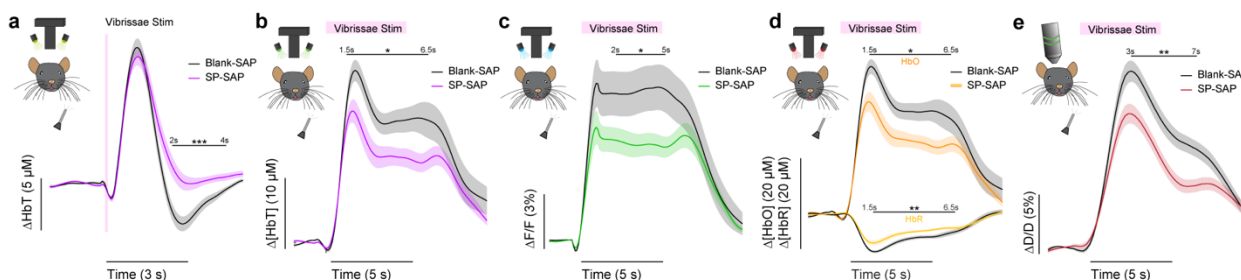


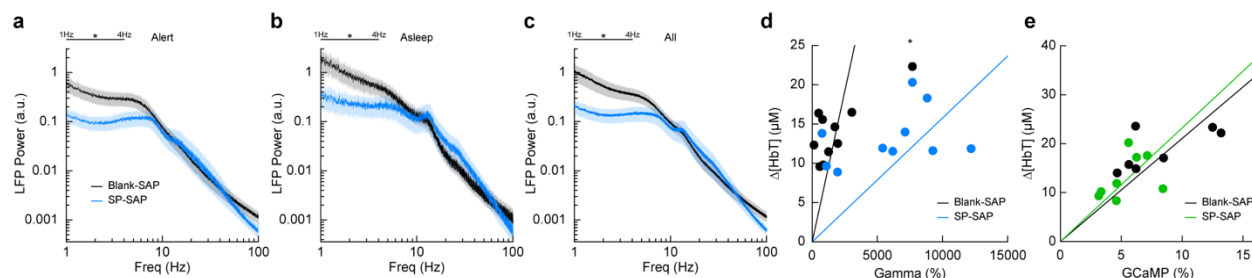
Figure 3. Ablation of type-I nNOS neurons reduces stimulus-evoked hemodynamic responses (a) Change in total hemoglobin in response to brief (0.1 second) vibrissae stimulation. (b) Change in total hemoglobin in response to extended (5 seconds) vibrissae stimulation. (c) Change in GCaMP fluorescence in response to extended vibrissae stimulation. (d) Change in oxy- and deoxy- hemoglobin in response to extended vibrissae stimulation. (e) Change in arteriole diameter in response to extended vibrissae stimulation. Error bars represent population averages \pm SEM. All statistics were evaluated between the indicated intervals. * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$, GLME.

182 following stimulus-onset decreased from $18.7 \pm 1.7 \mu\text{M}$ in Blank-SAP ($N = 7$, 3M/4F) to $13.1 \pm 1.4 \mu\text{M}$ in SP-SAP ($N = 8$, 4M/4F, GLME $p = 0.017$) (Fig. 3b). Calcium signals (Fig. 3c) evaluated 2.5 seconds following stimulus onset decreased from $8.2 \pm 1.3 \%$ in Blank-SAP to $5.3 \pm 0.7 \%$ in SP-SAP (GLME $p = 0.047$). $\Delta[\text{HbO}]$ (Fig. 3d) evaluated 1.5:6.5 seconds following stimulus onset decreased from $24.6 \pm 1.9 \mu\text{M}$ in Blank-SAP to $17.4 \pm 1.7 \mu\text{M}$ in SP-SAP (GLME $p = 0.010$) while $\Delta[\text{HbR}]$ evaluated from 1.5:6.5 seconds increased from $-6.0 \pm 0.4 \mu\text{M}$ in Blank-SAP to $-4.4 \pm 0.3 \mu\text{M}$ in SP-SAP (GLME $p = 0.005$). Evaluation windows were determined based on the duration of stimuli and the delayed onset of the hemodynamic response.

189 We further investigated single arterial dynamics during vibrissae stimulation with 2-photon microscopy. 190 Following ablation of type-I nNOS neurons, after vibrissae stimulation there was a decrease in arteriole diameter 191 (Fig. 3e) evaluated 3:7 seconds following stimulus onset from $11.6 \pm 0.8 \%$ in Blank-SAP ($N = 9$, 5M/4F, $n = 81$ 192 arterioles) down to $7.8 \pm 0.6 \%$ in SP-SAP ($N = 7$, 2M/5F, $n = 70$ arterioles, GLME $p = 0.008$). The arterial changes 193 closely mirror the $\Delta[\text{HbT}]$ changes, consistent with the substantial contribution of arterial dynamics to the blood 194 volume signal (Huo, 2015). We also evaluated changes in blood volume during voluntary locomotion, where we saw 195 a similar trend in decrease in locomotion-evoked $\Delta[\text{HbT}]$ evaluated 1.5:2.5 seconds following locomotion onset from 196 $17.5 \pm 2.0 \mu\text{M}$ in Blank-SAP ($N = 7$, 5M/2F) to $12.9 \pm 2.0 \mu\text{M}$ in SP-SAP ($N = 7$, 2M/5F), but it was not statistically 197 significant (GLME $p = 0.10$). These results show the removal of a very small number of neurons can drive substantial 198 reductions in both hemodynamic and neural response to sensory stimulation.

199 Type-I nNOS ablation reduces low frequency neural activity

200 To determine the effect that type-I nNOS neuron removal had on neural activity, we assessed the power 201 spectrum of the LFP in different arousal states, which we classified into alert, asleep, and all data (Turner, 2020). 202 Data was classified into the alert state when 15-minute blocks were predominantly (> 80%) in the awake state. Alert 203 periods contained fidgeting movements and bouts of whisking interspersed with awake quiescence. Periods 204 classified as asleep were predominantly composed of sleeping states (NREM/REM > 80% of all classifications in a 15- 205 minute block) with only brief periods of wakefulness, typically occurring during transitions between sleep states. The 206 'all' state includes everything irrespective of arousal state classification. The first ~60 minutes of each recording



207 **Figure 4. Type-I nNOS ablation alters low frequency neural activity and gamma-band neurovascular coupling** (a) Local field potential within the 208 vibrissae representation of somatosensory cortex during periods of Alert. (b) Asleep. (c) All data. (d) Change in gamma-band power vs. $\Delta[\text{HbT}]$ following 209 brief (0.1 seconds) vibrissae stimulation. (e) Change in GCaMP7s fluorescence vs. $\Delta[\text{HbT}]$ following prolonged (5 seconds) vibrissae 210 stimulation. Error bars represent population averages \pm SEM. All statistics were evaluated between the indicated intervals. * $p < 0.05$, ** $p < 0.01$, 211 *** $p < 0.001$, GLME (a-c), GLM (d, e).

207 session which included vibrissae stimulation was excluded. There was a pronounced reduction in the delta-band (1-
208 4 Hz) power of the LFP in all three arousal state categories. The power in the delta-band of the LFP in Blank-SAP mice
209 ($N = 9$, 4M/5F) in the alert state (**Fig. 4a**) was $3.4 \times 10^{-10} \pm 1.0 \times 10^{-10}$ a.u. compared to $1.0 \times 10^{-10} \pm 3.0 \times 10^{-11}$ a.u. in the
210 SP-SAP mice ($N = 9$, 5M/4F, GLME $p = 0.011$). The power in the delta-band of the LFP in Blank-SAP mice ($N = 7$, 3M/4F)
211 in the asleep state (**Fig. 4b**) was (NREM + REM) was $8.5 \times 10^{-10} \pm 3.0 \times 10^{-10}$ a.u. compared to $2.4 \times 10^{-10} \pm 9.1 \times 10^{-11}$ a.u.
212 in the SP-SAP mice ($N = 7$, 4M/3F, GLME $p = 0.022$). The power in the delta-band of the LFP in Blank-SAP mice ($N =$
213 9, 4M/5F) averaged across all arousal states (**Fig. 4c**) was $5.2 \times 10^{-10} \pm 1.7 \times 10^{-10}$ a.u. compared to $1.5 \times 10^{-10} \pm 4.3 \times 10^{-11}$
214 a. u. in the SP-SAP mice ($N = 9$, 5M/4F, GLME $p = 0.016$). These results parallel those seen in studies where knockout
215 of nNOS produces lower delta band power (Morality, 2013) and optogenetic stimulation of nNOS neurons produces
216 low frequency oscillations (Zielinski, 2019), suggesting that type-I nNOS neurons promote these slow oscillations via
217 NO release. In contrast to the observed reductions in LFP, we noted no gross changes in the power spectra of cerebral
218 blood volume fluctuations between groups during any arousal state (**Fig. S4**).

219 We next wanted to see whether the amount of neural activity and corresponding $\Delta[\text{HbT}]$ in response to vibrissae
220 stimulation was altered following type-I nNOS neuron ablation. As a measure of neurovascular coupling, we used
221 the slope of the line fitting the change in gamma-band power vs. $\Delta[\text{HbT}]$ following brief (0.1 second) vibrissae
222 stimulation (**Fig. 4d**). Decreases in the slope indicate a smaller vascular response for a given amount of neural
223 activity, indicating a decrease in neurovascular coupling. We found the slope was significantly increased in SP-SAP
224 mice compared to Blank-SAP mice (Blank-SAP: 0.02 ± 0.03 ; SP: 0.005 ± 0.006 , GLM $p = 0.0001$). We also evaluated
225 the change in GCaMP7s fluorescence vs. $\Delta[\text{HbT}]$ during prolonged (5 seconds) vibrissae stimulation from 2:5 seconds
226 (**Fig. 4e**) and saw no significant change in the slope (Blank-SAP: 251.3 ± 75.3 ; SP: 256.5 ± 73.2 , GLM $p = 0.34$). Given
227 that the sensory-evoked LFP is driven more by synaptic input, while GCaMP7s signals are driven by local neural
228 activity, these results indicate that the loss of type-I nNOS neurons reduces overall excitability, but that the input
229 drive from other areas might be increased, potentially due to homeostatic mechanisms at the input synapses
230 (Turrigiano, 2008).

231 **Neurovascular coupling was weakly affected by type-I nNOS removal**

232 We next looked at the cross-correlation between blood volume and neural signals, which provides a measure
233 of spontaneous neurovascular coupling. For electrophysiological measures, we measured the cross-correlation
234 between gamma-band power and blood volume (**Fig. 5a**). We evaluated the peak cross-correlation during the
235 resting-state, during long periods while alert, or long periods while asleep. The cross-correlation between neural
236 activity and hemodynamic signals is substantially higher during periods of behavior (whisking, fidgeting) than during
237 the resting-state because self-generated motion and whisking drive increases in neural activity and vasodilation
238 (Winder, 2017; Drew, 2019b; Stringer, 2019; Claron, 2023; Tu, 2024). The correlation between neural activity and
239 blood volume changes is higher during sleep than in the awake state (Turner, 2020). We saw no significant difference
240 in the peak cross-correlation during the resting-state (**Fig. 5b**) between Blank-SAP mice ($N = 9$, 4M/5F, 0.08 ± 0.007)

241 and SP-SAP mice ($N = 9$, 5M/4F, 0.06 ± 0.008 , GLME $p = 0.22$). We did observe a significant difference in the peak
 242 cross-correlation during both periods of alert and asleep, from 0.23 ± 0.01 (Blank-SAP) to 0.16 ± 0.03 (SP-SAP, GLME
 243 $p = 0.036$) during alert periods (**Fig. 5c**) and 0.40 ± 0.02 (Blank-SAP) dropping to 0.33 ± 0.02 (SP-SAP, GLME $p = 0.012$)
 244 while asleep (**Fig. 5d**). We next wanted to evaluate whether this drop in neurovascular coupling was more prevalent
 245 at a particular modulation frequency. When analyzing the coherence between the gamma-band power and
 246 hemodynamic signals during these different arousal states, we noted no dominant frequency and saw no significant
 247 changes in the lower frequencies
 248 associated with neurovascular coupling
 249 (≤ 0.5 Hz). During the resting-state (**Fig.**
 250 **5e**), the average coherence between 0.1
 251 and 0.5 Hz was 0.28 ± 0.008 in Blank-SAP
 252 mice and 0.23 ± 0.01 in SP-SAP mice
 253 (GLME $p = 0.23$). The average coherence
 254 between 0.01 and 0.5 Hz during alert
 255 periods (**Fig. 5f**) was 0.29 ± 0.004 in
 256 Blank-SAP mice and 0.25 ± 0.005 in SP-
 257 SAP mice (GLME $p = 0.06$), and during
 258 asleep periods (**Fig. 5g**) it was $0.34 \pm$
 259 0.008 in Blank-SAP mice and $0.32 \pm$
 260 0.007 in SP-SAP mice (GLME $p = 0.46$).
 261 Lastly, to evaluate any changes in the
 262 predictive power of the neural-
 263 hemodynamic relationship, we fit a
 264 hemodynamic response function using
 265 the gamma-band power and
 266 hemodynamic response following
 267 periods of vibrissae stimulation. We
 268 observed that ablation of Type-I nNOS
 269 neurons did not alter the predictive
 270 power of the hemodynamic response
 271 function (**Fig. S5**).
 272 We next evaluated neurovascular
 273 coupling from optical measures of bulk
 274 activity using GCaMP7s (**Fig. 5h**). The
 275 peak resting-state cross-correlation (**Fig.**

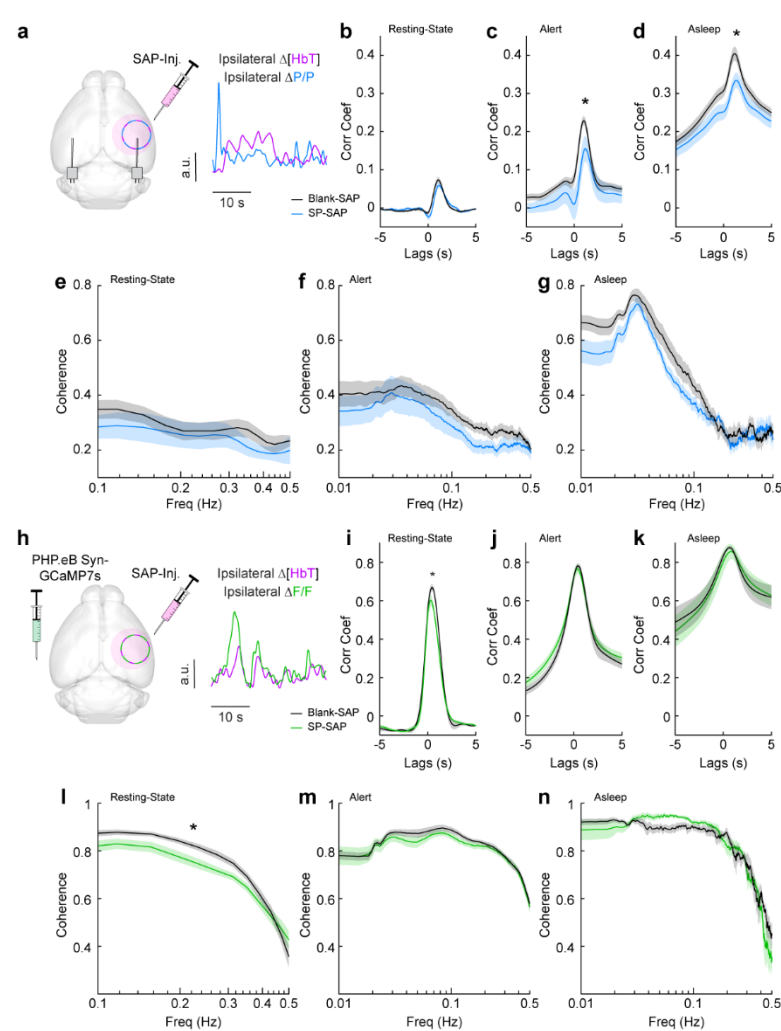


Figure 5. Neurovascular coupling was only weakly affected by type-I nNOS removal (a) Schematic demonstrating intracortical injection of either Blank-SAP or SP-SAP and the analysis of gamma-band power and hemodynamic signals from within the vibrissae representation of somatosensory cortex, $N = 9$ mice per group. (b) Gamma- Δ [HbT] resting-state cross-correlation (c) Gamma- Δ [HbT] alert cross-correlation (d) Gamma- Δ [HbT] asleep cross-correlation (e) Gamma- Δ [HbT] resting-state coherence (f) Gamma- Δ [HbT] alert coherence (g) Gamma- Δ [HbT] asleep coherence (h) Schematic demonstrating intracortical injection of either Blank-SAP or SP-SAP and the analysis of GCaMP7s fluorescence and hemodynamic signals from within the vibrissae cortex, $n = 6-7$ mice per group. (i) GCaMP7s- Δ [HbT] resting-state cross-correlation (j) GCaMP7s- Δ [HbT] alert cross-correlation (k) GCaMP7s- Δ [HbT] asleep cross-correlation (l) GCaMP7s- Δ [HbT] resting-state coherence (m) GCaMP7s- Δ [HbT] alert cross-correlation (n) GCaMP7s- Δ [HbT] asleep cross-correlation. Error bars represent population averages \pm SEM. * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$, GLME.

276 **5i)** between GCaMP7s fluorescence and ongoing hemodynamics showed a significant decrease, 0.68 ± 0.02 with
277 Blank-SAP versus 0.62 ± 0.02 with SP-SAP (GLME $p = 0.038$). However, there was no drop in peak cross-correlation
278 during either alert or asleep as seen with the gamma-band power. During alert periods (**Fig. 5j**), the peak GCaMP7s
279 cross-correlation was 0.78 ± 0.12 with Blank-SAP and 0.77 ± 0.02 with SP-SAP (GLME $p = 0.54$) while during asleep
280 periods (**Fig. 5k**) it was 0.88 ± 0.008 with Blank-SAP and 0.86 ± 0.04 with SP-SAP (GLME $p = 0.58$). Like the peak in
281 cross-correlation, the average resting-state coherence (**Fig. 5l**) did show a significant drop across the lower
282 frequencies, 0.75 ± 0.01 with Blank-SAP vs. 0.70 ± 0.01 with SP-SAP (GLME $p = 0.049$). There was no drop in coherence
283 during alert periods (**Fig. 5m**) at 0.78 ± 0.005 with Blank-SAP and 0.77 ± 0.004 with SP-SAP (GLME $p = 0.42$) nor during
284 asleep periods (**Fig. 5n**) at 0.75 ± 0.009 with Blank-SAP and 0.74 ± 0.01 with SP-SAP (GLME $p = 0.63$). While there are
285 small changes in neural-hemodynamic correlations, the differences in neurovascular coupling across arousal states
286 were relatively small following localized removal of type-I nNOS neurons, meaning any effects of ablation on
287 hemodynamic responses are primarily mediated by changes in neural activity.

288 **Type-I nNOS ablation reduces low-frequency interhemispheric coherence**

289 Because type-I nNOS neurons send and receive many modulatory signals across a range of distances, they could
290 help serve to coordinate neural and vascular dynamics across the brain. We tested this hypothesis by comparing
291 how loss of type-I nNOS neurons changed the coherences of neural and hemodynamics signals between the left and
292 right vibrissae cortex, which are generally highly correlated across all behaviors and frequencies (Turner, 2020).
293 Neural activity is also bilaterally correlated, though less so than vascular dynamics. Average resting-state coherence
294 between the left and right $\Delta[\text{HbT}]$ signals (**Fig. 6a**) in the vibrissae cortex were 0.79 ± 0.005 for Blank-SAP and $0.73 \pm$
295 0.008 for SP-SAP mice ($p = 0.014$, GLME, **Fig. 6b**). This reduction in coherence was seen over all frequencies (0-0.5
296 Hz) during both alert periods (**Fig. 6c**), at 0.86 ± 0.002 for Blank-SAP and 0.81 ± 0.003 for SP-SAP ($p = 7.4 \times 10^{-6}$, GLME),
297 as well as asleep periods (**Fig. 6d**) with 0.86 ± 0.003 for Blank-SAP and 0.79 ± 0.005 for SP-SAP ($p = 0.0008$, GLME).
298 We then looked at the coherence of gamma-band power across hemispheres, where average coherence for bilateral
299 gamma-band signals in the resting-state were 0.19 ± 0.006 (Blank-SAP) and 0.19 ± 0.01 (SP-SAP, $p = 0.88$, GLME, **Fig**
300 **6e, f**). There was no significant difference during the alert periods (**Fig 6g**) at 0.22 ± 0.005 (Blank-SAP) and $0.20 \pm$
301 0.006 (SP-SAP, $p = 0.43$, GLME), but ablation of type-I nNOS neuros produced a reduction of coherence during asleep
302 periods (**Fig. 6h**) at 0.35 ± 0.009 (Blank-SAP) and 0.29 ± 0.009 (SP-SAP, $p = 0.047$, GLME).

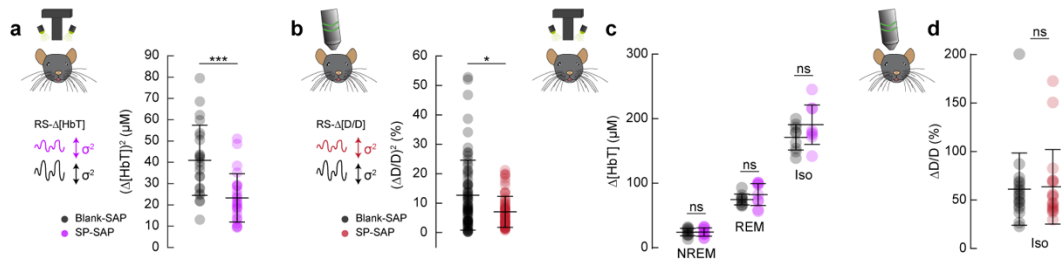
303 Average resting-state coherence across bilateral $\Delta[\text{HbT}]$ signals (**Fig. 6i**) taken from the vibrissae cortex were
304 0.83 ± 0.005 (Blank-SAP) and 0.78 ± 0.007 (SP-SAP, $p = 0.021$, GLME, **Fig. 6j**). This reduction in coherence persisted
305 across all frequencies during the both alert periods (**Fig. 6k**) at 0.92 ± 0.001 (Blank-SAP) and 0.90 ± 0.001 (SP-SAP, p
306 $= 0.012$, GLME) as well as sleep periods (**Fig. 6l**) at 0.90 ± 0.002 (Blank-SAP) and 0.83 ± 0.005 (SP-SAP, $p = 0.004$,
307 GLME). Average resting-state coherence across bilateral GCaMP7s signals (**Fig. 6m**) taken from the vibrissae cortex
308 were 0.75 ± 0.009 (Blank-SAP) and 0.70 ± 0.009 (SP-SAP, GLME $p = 0.12$, **Fig. 6n**). This reduction in coherence
309 persisted across all frequencies during the both alert periods (**Fig. 6o**) at 0.88 ± 0.003 (Blank-SAP) and 0.84 ± 0.003

310 (SP-SAP, $p = 0.029$, GLME) but not in sleep periods (Fig. 6p) at 0.85 ± 0.005 (Blank-SAP) and 0.84 ± 0.005 (SP-SAP, GLME $p = 0.52$). When the correlation of bilateral signals was evaluated with Pearson's correlation coefficients, the same general trend remained (Fig. S6). These results show that removal of type-I nNOS neurons reduces both 313 vascular and neural coordination across hemispheres.

314 **Type-I nNOS neurons control vasomotion amplitude but not baseline diameter**

315 We next wanted to establish how ablation of type-I nNOS neurons affected vasomotion, spontaneous 316 oscillations in the absence of behavior, measured at the scale of blood volume and at the level of single arterioles, 317 as well as the basal tone of blood vessels. The variance in $\Delta[\text{HbT}]$ during rest, a measure of vasomotion amplitude, 318 was significantly reduced following type-I nNOS ablation (Fig. 7a), dropping from $40.9 \pm 3.4 \mu\text{M}^2$ in the Blank-SAP 319 group ($N = 24$, 12M/12F) to $23.3 \pm 2.3 \mu\text{M}^2$ in the SP-SAP group ($N = 24$, 11M/13F) (GLME $p = 6.9 \times 10^{-5}$). Individual 320 pial and penetrating arterioles showed the same reduction in vasomotion after type-I nNOS neuron ablation, with 321 Blank-SAP ($N = 9$, 5M/4F, $n = 70$ arterioles) having a resting diameter variance of $12.6 \pm 1.4 \%^2$ and SP-SAP ($N = 7$, 322 2M/5F, $n = 65$ arterioles) of $8.0 \pm 0.8 \%^2$ (Fig. 7b).

323 If type-I nNOS neurons control the basal diameter of vessels via secreted vasodilators, potentially in an arousal



324 **Figure 7. Type-I nNOS ablation reduces resting-state hemodynamics and low frequency neural activity** (a) Variance in resting-state 325 hemodynamics signals measured with widefield optical imaging. (b) Variance in resting-state arteriole diameter measured with two-photon. (c) 326 Average $\Delta[\text{HbT}]$ during periods of NREM sleep, periods of REM sleep, and following administration of isoflurane ($n = 9$ mice per group). (d) Change 327 in normalized arteriole diameter following administration of isoflurane. Error bars represent population averages \pm SD. All statistics were 328 evaluated between the indicated intervals. * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$ GLME.

329 state dependent way, when we ablate these neurons we would observe a difference in blood volume across arousal 330 states and in the maximal dilation elicited with isoflurane (Gao, 2015). We saw no changes in the average change in 331 $\Delta[\text{HbT}]$ across states (Fig. 7c, Fig. S7). The resting-state $\Delta[\text{HbT}]$ in Blank-SAP mice was $0.29 \pm 0.7 \mu\text{M}$ compared to 332 $0.29 \pm 0.9 \mu\text{M}$ in SP-SAP (GLME $p = 0.997$). NREM $\Delta[\text{HbT}]$ in Blank-SAP mice was $23.9 \pm 5.8 \mu\text{M}$ compared to $24.1 \pm$ 333 $6.5 \mu\text{M}$ in SP-SAP (GLME $p = 0.96$). REM $\Delta[\text{HbT}]$ in Blank-SAP mice was $74.5 \pm 8.4 \mu\text{M}$ compared to $82.3 \pm 17 \mu\text{M}$ in 334 SP-SAP (GLME $p = 0.26$). The $\Delta[\text{HbT}]$ following isoflurane in Blank-SAP mice was $170.9 \pm 19.7 \mu\text{M}$ compared to $190.5 \pm 30.5 \mu\text{M}$ in SP-SAP (GLME $p = 0.15$). We repeated the isoflurane experiment on a set of pial and penetrating 335 arterioles under 2-photon microscopy. The change in normalized arteriole diameter following administration of 336 isoflurane ($n = 7-9$ mice per group, 18-19 arterioles per group) was $61 \pm 37.2\%$ in Blank-SAP compared with $63.6 \pm$ 337 38.4% in SP-SAP ($p = 0.85$, ttest). These results indicate that type-I nNOS neurons play a role in driving spontaneous 338 hemodynamic fluctuations, but not in setting the basal diameter during different states.

335 **Discussion**

336 We selectively ablated type-I nNOS neurons from the somatosensory cortex, which had marked effects on
337 neural activity and vascular dynamics, but minimal changes in neurovascular coupling. These results are surprising
338 given previous work showing stimulation of these neurons in isolation causes vasodilation and minimal neural
339 activity changes. Our approach of using SP-conjugated saporin allowed a nongenetic means of targeting a critical
340 neuronal cell type, supporting further exploration of the role of type-I nNOS neurons in transgenic mouse models of
341 disease without complicated breeding schemes, as well as in non-model organisms. Our results point to these
342 neurons being an important orchestrator of neural and vascular dynamics, as loss of these neurons causes
343 desynchronization between hemispheres of both neural and vascular signals, as well as altered responses to sensory
344 stimulation. While previous studies using specific activation of type-I nNOS neurons have emphasized the roles of
345 these neurons in driving vasodilation with minimal changes in neural activity, our results point to a larger role of
346 these neurons in organizing and patterning both spontaneous and sensory-evoked neural activity. This is likely due
347 to a loss in NO signaling as well as a loss of the many of the other neuropeptides expressed by type-I nNOS neurons,
348 which all have known effects on other neurons.

349 While previous studies have found that activation of type-I nNOS neurons, either optogenetically or via
350 administration of substance P, results in vasodilation (which would imply an important role for these neurons for
351 neurovascular coupling), our work provides insight into the integrated function of type-I nNOS neurons in neural
352 circuits and vascular dynamics. Importantly, methodological approaches differentiate our strategy from previous
353 published work. One interpretation is that there could be compensatory rewiring of local and/or network activity
354 following type-I nNOS ablation, where other signaling pathways from the neurons to the vasculature become
355 strengthened to compensate for the loss of vasodilatory signaling from the type-I nNOS neurons. While this likely
356 happens to some degree, this interpretation is less likely to completely account for the effect given these potential
357 compensatory changes do not prevent large changes in neural activity and resting vasomotion. Second, there may
358 be some non-linear interactions between vasodilatory signals and/or the vasodilator mechanisms that are arousal-
359 state-dependent. The diameter of arteries tracks the smooth muscle membrane potential linearly, up until a
360 saturation point above which any hyperpolarization does not induce further dilation (Knot, 1998; Wölflé, 2011; Hill,
361 2012). Many vasoactive pathways from neurons to vessels are known to exist, whose additive individual effects could
362 account for larger than biologically possible changes in vasodilation (Hosford, 2019). It may be that for dilation
363 occurring during sensory stimulation and sleep, the vasodilatory stimulus to vessels exceeds the saturation point, so
364 that the loss of one vasodilatory pathway does not further affect the vascular response (**Fig. S8**). This would also be
365 consistent with isolated optogenetic or chemical stimulation of type-I nNOS neurons being able to drive substantial
366 dilation, while the loss of these neurons does not have a large impact on the vascular response.

367 Finally, pharmacological ablation and optogenetic/chemogenetic activation are not mirror manipulations and
368 have differences that can produce non-symmetrical changes. Symmetrical changes would only occur if the neural
369 circuitry and the signaling to the vessel were completely linear. Optogenetic activation/deactivation of a single cell

370 type does not produce symmetrical changes in the activity of other neurons (Phillips, 2016), so there is little reason
371 to think ablation would have the exact opposite effect as activation. Furthermore, optogenetic/pharmacological
372 activation of a single cell type is unlikely to occur endogenously, as neural activity across different cell types is largely
373 correlated (Bugeon, 2022). For example, during whisker stimulation, not only are type-I nNOS neurons in the
374 somatosensory cortex activated (Ruff, 2024), but nearly every other cell type is as well (Staiger, 2021), which mean
375 that type-I nNOS neurons are not the only neurons, and not even the only NO producing neurons, sending signals to
376 the vasculature during sensory stimulation.

377 One surprise is that we observed no changes in the vasodilation during NREM sleep, where type-I nNOS neurons
378 are known to be active and play a role in inducing sleep (Kilduff, 2011). During NREM sleep, there is marked arterial
379 dilation (Turner, 2020; Gheres, 2023), and it would seem natural that type-I nNOS neurons might drive this. However,
380 we saw no difference in NREM dilation between the ablated and control mice. Again, this could be due to
381 compensation, saturation of the vasodilatory response, or it could be due to type-I nNOS neurons releasing other
382 vasoconstrictory peptides (SST, NPY) so their net effects are cancelled out. This last possibility is consistent with the
383 loss of the post-stimulus vasoconstriction seen after type-I nNOS neuron ablation. Again, like with sensory
384 stimulation, there may be a non-linearity in the vascular responses, so that another large factor, such as large
385 neuromodulatory changes during sleep, could dominate over any other vascular signaling factor. The large
386 norepinephrine decreases during NREM (Osorio-Forero, 2021; Kjaerby, 2022; Turner, 2023) could release the vessels
387 from tonic vasoconstriction (as norepinephrine is a vasoconstrictor (Bekar, 2012)), resulting in the dilation during
388 NREM.

389 Finally, we saw a marked reduction in spontaneous vascular oscillation (vasomotion) at rest, both at the level of
390 single arteries, and at the level of blood volume. These spontaneous oscillations move cerebrospinal fluid (CSF) (van
391 Veluw, 2020; Kedarasetti, 2022; Holstein-Ronsbo, 2023) which is important for clearing waste from the brain, and a
392 reduction in amplitude will decrease the pumping efficacy. Additionally, there is a loss of coordination of neural and
393 vascular dynamics across hemispheres after ablation of type-I nNOS neurons, indicated by the drop in coherence.
394 Although a loss of synchronous dilations/constrictions might affect CSF movement, we might imagine that it would
395 also adversely impact CSF pumping. Aged human brains show reduced interhemispheric synchrony in resting-state
396 signals (Zhao, 2020), analogous to what we see here. Type-I nNOS neurons also seem uniquely vulnerable to stress
397 (Han, 2019), and loss of type-I nNOS neurons reduces power in the 1-4Hz band of the LFP, which is positively
398 associated with CSF clearance (Hablitz, 2019; Jiang-Xie, 2024). It is possible that adverse life experiences could cause
399 the loss of type-I nNOS neurons, leading to reduction in CSF flow seen in age (Kress, 2014; Matrongolo, 2023) that is
400 thought to correlate with neurodegeneration.

401 **Materials and Methods**

402 This study was performed in accordance with the recommendations of the Guide for the Care and Use of
403 Laboratory Animals of the National Institutes of Health. All procedures were performed in accordance with protocols

404 approved by the Institutional Animal Care and Use Committee of Pennsylvania State University (Protocol
405 201042827). Data were acquired from 119 C57BL6/J mice (#000664, The Jackson Laboratory, Bar Harbor, ME)
406 compromised of 57 males and 62 females between 3 and 9 months of age. Food and water were provided *ad libitum*
407 and animals were housed on a 12-hr. light/dark cycle with all experiments occurring during the light cycle. Mice were
408 individually housed after surgery. Sample sizes are consistent with previous studies (Winder, 2017; Turner, 2020;
409 Turner, 2023) and experimenters were not blind to experimental conditions or data analysis except for histological
410 experiments.

411 **Surgical Procedures.** Saporin is a ribosome-inactivating protein that was conjugated to the Sar⁹, Met(O₂)¹¹ analog of
412 Substance P (SP-SAP) or to a control peptide (Blank-SAP) (IT11 & IT21, Advanced Targeting Systems, Carlsbad, CA).
413 Mice were anesthetized using 5% isoflurane (2% maintenance) vaporized in pure oxygen and were then injected
414 intracortically with 4 ng of either SAP conjugate or Blank-SAP in 100 nL of artificial cerebrospinal fluid (aCSF). The
415 incision site was sterilized with betadine and 70% ethanol followed by a retraction of the skin atop the skull. A small
416 (< 0.5 mm) craniotomy was made above the vibrissae representation of somatosensory cortex (2 mm caudal, 3.25
417 mm lateral) and a sterile glass-pulled needle (tip diameter 50-100 μ m) was inserted 500 μ m beneath the cortical
418 surface at 45°. The SAP conjugate was slowly injected at 100 nL/min using a programmable syringe pump (Harvard
419 Apparatus, Holliston, MA) followed by closure of the incision with VetBond (3M, Saint Paul, MN). A subset of animals
420 were also injected retro-orbitally with 25 μ L of AAV PHP.eB-syn-jGCaMP7s-WPRE [2x10¹³ GC/mL] (104487-PHPeB,
421 Addgene, Watertown, MA) diluted in 25 μ L of sterile saline. Animals were given at least 2 weeks to recover prior to
422 undergoing additional procedures. For imaging, a custom-machined titanium head bar was adhered atop the
423 occipital bone of the skull using cyanoacrylate glue (Vibra-Tite 32402, ND Industries, Clawson, MI) and dental cement
424 (Ortho-Jet, Lang Dental, Wheeling, IL). Self-tapping 3/32' #000 screws (J.I. Morris, Oxford, MA) were implanted in
425 the frontal bones for structural stability. Electrodes were implanted into cortex and hippocampus using PFA-coated
426 tungsten stereotrodes (#795500, AM systems, Sequim, WA) for recordings of local field potentials (LFP) and into the
427 neck muscles using a pair of PFA-coated 7-strand stainless-steel wires (#793200, AM systems, Sequim, WA) for
428 electromyography (EMG). Polished and reinforced thinned-skull windows (Drew et al., 2010a; Shih et al., 2012b)
429 were implanted over the somatosensory areas (either bilaterally or right hemisphere) using #0 coverslips (#72198,
430 Electron Microscopy Sciences, Hatfield, PA). Detailed surgical procedures are as previously described (Turner, 2020).

431 **Data acquisition.** Data were acquired with a custom LabVIEW program (National Instruments;
432 <https://github.com/DrewLab/LabVIEW-DAQ>). Details on widefield optical imaging, electromyography (EMG),
433 electrophysiology, vibrissae stimulation, and behavioral measurements including tracking of vibrissae and pupil
434 diameter were performed as previously described (Turner, 2020; Zhang, 2022; Turner, 2023)). Mice were gradually
435 acclimated to head-fixation of increasing duration (15, 30, 60 minutes) on the days preceding the onset of
436 experiments. The vibrissae (left, right, or a third air puffer not directed at the body as an auditory control) were
437 randomly stimulated with air puffs [0.1 seconds or a train of 5 second pulses, 10 pounds force per square inch (PSI)]

438 occurring every 30–45 seconds for the first 1 hour of imaging. Data were acquired in 15-minute intervals with a brief
439 (~30 seconds) gap in between for saving data to disk. Each animal was run for up to 6 imaging sessions lasting from
440 1–5 hours depending on experiment.

441 **Histology.** Following the conclusion of imaging experiments, animals were deeply anesthetized and transcardially
442 perfused with heparin-saline followed by 4% paraformaldehyde. Presence or absence of nNOS-positive neurons in
443 the injected hemisphere was verified using nicotinamide adenine dinucleotide phosphate (NADPH)-diaphorase
444 staining for localizing the sparsely populated type-I nNOS neurons (Scherer-Singler, 1983).

445 **Immunohistochemistry.** All histological analyses were done blinded to the experimental condition. Mice were
446 deeply anesthetized with 5% isoflurane and perfused transcardially with ice-cold phosphate buffered saline (PBS, pH
447 7.4) and 4% paraformaldehyde (PFA, pH 7.4). Brains were removed, post-fixed in PFA for 24 hours and stored in PBS
448 at 4°C for less than 1 week. A fiduciary mark was placed in the right hemisphere. 40-μm free floating sections were
449 sliced with a Leica vibratome (VS 1200, Leica) and stored in PBS for less than 1 week. Prior to immunostaining, slices
450 were washed three times in PBS for 10 minutes each, and underwent antigen retrieval in 10 mM sodium citrate
451 buffer (pH 6.0) at 80°C for 30 minutes. Slices were washed three times in PBS for 10 minutes each, and permeabilized
452 in 0.5% Triton X-100 in PBS for 60 minutes. Nonspecific binding was blocked with 5% normal donkey serum (NDS)
453 (ab7475) in 0.1% Triton X-100 in PBS for 60 minutes. Slices were then incubated in a primary antibody cocktail,
454 including goat anti-nNOS (1:500, ab1376), rabbit anti-IBA-1 (1:500, ab178847), rat anti-GFAP (1:500, ThermoFisher
455 13-0300), rabbit anti NeuN (1:2000, EnCor), rabbit anti-TACR1 (1:500 Invitrogen PA1-16713) in 2.5% NDS in 0.1%
456 Triton X-100 in PBS for 48-h at 4°C. Slices were rinsed three times with PBS for 10 minutes each, and incubated in a
457 fluorophore-tagged secondary antibody cocktail, including donkey anti-rabbit Alexa Fluor 488 (1:500, ab150073),
458 donkey anti-goat Alexa Fluor 594 (1:500, ab150132), donkey anti rabbit Alexa Fluor 647 (1:500, ab150075), donkey
459 anti rat Alexa Fluor 488 (1:500 ab150153), for 4 hours at room temperature. Slices were rinsed again three times
460 with PBS, with the last step including DAPI (1:10,000, 10mg/mL, Millipore Sigma, 10236276001), mounted on glass
461 slides, air-dried and cover slipped with Immunomount (Thermo Fisher Scientific, Waltham, MA, United States).
462 Images were obtained with an Olympus BX63 upright microscope (Center Valley, PA, United States) under matched
463 exposure settings. Three to eight images from both hemispheres were taken per region.

464 **Cell counting and immunofluorescence quantification.** Total cell counts and absolute changes in
465 immunofluorescence (Fig. 1) were quantified using ImageJ (National Institutes of Health, Bethesda, MD, United
466 States). For total cell counts, a region of interest (ROI) was delineated, and cells were automatically quantified under
467 matched criteria for size, circularity and intensity. Each ROI's total cell count was divided by the ROI's area to give a
468 total density value (Dao, 2020; Smith, 2020; Sicher, 2023). Immunoreactivity was quantified as mean fluorescence
469 intensity of the ROI (Pleil, 2015).

470

471 **Data Analysis.** Data were analyzed with code written by K.L.T, M.S.H, Q.Z, K.W.G, and P.J.D. (MATLAB 2019b–2024a,
472 MathWorks).

473 **Statistical analysis.** Statistical evaluations were made using either generalized linear mixed effects (GLME), unpaired
474 t-test, or general linear model (GLM). GLME models had the arousal state as a fixed effect, mouse identity as a
475 random effect, and hemisphere [left/right (L/R), if applicable] as an interaction with the animal ID, or using a paired
476 t test where appropriate. Unless otherwise stated, statistical results report p values from a GLME test. All reported
477 quantifications are mean \pm SD unless otherwise indicated. Unless otherwise noted, all pupil diameter measurements
478 are in z-units. MATLAB functions used were fitglme, ttest, fitglm.

479 **Hemodynamic correction.** Widefield imaging was done with a Dalsa 1M60 Pantera CCD camera (Phase One,
480 Cambridge, MA) with a magnifying lens (VZM 300i, Edmund Optics, Barrington, NJ). Reflectance measurements (Fig.
481 2) were converted to changes in total hemoglobin ($\Delta[\text{HbT}]$), oxy-hemoglobin ($\Delta[\text{HbO}]$), and deoxy-hemoglobin
482 ($\Delta[\text{HbR}]$) using the Beer–Lambert law (Ma et al., 2016a, b). Correction for attenuation of GCaMP7s fluorescence due
483 to absorption of the surrounding tissue was corrected as previously described (Kramer, 1979; Ma, 2016b; Wright,
484 2017; Scott, 2018). Changes in fluorescence intensity in the GCaMP7s signal due to blood absorption were
485 approximated in a pixel-wise fashion by multiplying each value by the ratio of the green and blue channel's resting
486 baseline pixel value.

487 **Electrophysiology.** Gamma-band [30–100Hz] LFP band was digitally bandpass filtered from recorded broadband
488 data using a third-order Butterworth filter, squared, low-pass filtered below 1 Hz, and resampled at 30 Hz. Time-
489 frequency spectrograms (Fig. 2) were calculated using the Chronux toolbox version 2.12 v03 (Bokil, 2010), function
490 mtspecgramc with a 5 s window and 1/5 s step size using [5,9] tapers and a passband of 1–100 Hz to encompass the
491 LFP. EMG (300–3 kHz) from the neck muscles was bandpass filtered, squared, convolved with a Gaussian kernel with
492 a standard deviation of 0.5 seconds, log transformed, then resampled at 30 Hz. MATLAB function(s): butter, zp2sos,
493 filtfilt, gausswin, log10, conv, resample.

494 **Evoked responses and slope.** Evoked responses (Fig. 3) including whisker stimulation and locomotion for the various
495 data types ([HbT], [HbO], [HbR], GCaMP7s, arteriole diameter) were compared between the indicated intervals (i.e.,
496 2:4 seconds post-stimulation). The mean of the 2 seconds preceding event onsets were subtracted from the event
497 and smoothed with a 3rd order Savitzky-Golay filter. The average slope was calculated by comparing the rise ($\Delta[\text{HbT}]$)
498 over the run (neural data) before being fit with a linear model forced through the origin. MATLAB function(s):
499 sgolayfilt)

500 **Spectral power and coherence.** Spectral power (Fig. 4, 5, 6 S5) was estimated using the Chronux toolbox (Bokil,
501 2010) function mtspectrumc. Data was detrended within individual events and truncated to the desired length

502 depending on behavior (10 seconds for rest, 15 minutes for Alert and Asleep). Coherence analysis between two
503 signals within the same hemisphere or between signals recorded bilaterally was run for each data type Chronux
504 function coherencyc after detrending using the MATLAB function detrend.

505 **Cross-correlation.** Cross-correlations (Fig. 5) between gamma-band power or $\Delta F/F$ and changes in total hemoglobin
506 $\Delta[HbT]$ were taken during periods of resting-state, alert, and asleep. Data were mean-subtracted and digitally
507 lowpass filtered (< 1 Hz) with a fourth-order Butterworth filter (MATLAB function(s): butter, zp2sos, filtfilt). Cross-
508 correlation analysis was run for each arousal state (MATLAB function(s): xcorr) with a ± 5 second lag time.

509 **Resting-state variance and $\Delta[HbT]$ during different arousal states.** Variance during the resting-state for both $\Delta[HbT]$
510 and diameter signals (Fig. 7) was taken from resting-state events lasting ≥ 10 seconds in duration. Average $\Delta[HbT]$ in
511 each arousal state was taken with respect to awake resting baseline events ≥ 10 seconds in duration. Continuous
512 NREM sleep events ≥ 30 seconds, REM sleep events ≥ 60 seconds, and periods following administration of isoflurane
513 were compared between groups. Each event was digitally lowpass filtered (<1 Hz) with a fourth-order Butterworth
514 filter (MATLAB function(s): butter, zp2sos, filtfilt) and then averaged within each individual time series prior to
515 comparing across animals/groups.

516

517 **Open field behavior.** Exploratory behavior (Fig. S2) was measured in a custom-made 30 cm x 60 cm arena during
518 the light phase. Mice were not habituated to the arena. Mice were placed in the arena for 10 minutes to explore,
519 and their behavior was recorded using an overhead camera (Blackfly BFLY-U3-23S6M, Teledyne FLIR, Wilsonville,
520 OR) with 6 mm fixed focal length lens (Edmund Optics, Barrington, NJ, Stock #33939) at a frame rate of either 15 or
521 30 frames per second. The arena was illuminated with 780nm light. The surroundings of the arena were dark during
522 the entire recording session. The arena was cleaned with 70% ethanol in between animals. Only the first 5 minutes
523 of the behavior were analyzed and reported here. The total distance traveled over the first 5 minutes and the time
524 spent in the 25 x 55 cm center rectangle were quantified. Generalized linear mixed effect (GLME) was used in
525 MATLAB to perform statistical analysis. No statistical outliers were removed from the data. Mouse behavior was
526 tracked using DeepLabCut (Mathis, 2018) and analyzed with custom MATLAB algorithms (Zhang, 2022; Brockway,
527 2023). Eight points on the body (left ear, right ear, head, mid, back near the hip joint, base of the tail, midpoint of
528 the tail, and end of the tail) were tracked. Four markers placed in the arena's four corners with an additional four
529 points calculated from the four corners to track the center rectangle. A DeepLabCut model was trained and
530 evaluated on a subset of mice before applying the model to all the mice. DeepLabCut tracking was considered
531 acceptable if the tracking confidence was above 97%; however, in most cases, it was higher than 99%. Tracking
532 positions were exported to a CSV file containing the tracked location's XY coordinates (frame pixels). Tracked videos
533 from random mice were visualized to confirm the accuracy of the tracking. The experimenter was blinded to drug
534 injection group identification until tracking and analysis were completed. The distance was calculated from the first
535 5 minutes as Euclidean distance between the point tracked in the middle of the mouse body from subsequent

536 frames. Center time was quantified as the number of frames a mouse spends within the polygon bounded by 4 points
537 tracked on the center of the arena.

538 **Sleep scoring.** Sleep scoring (**Fig. S3**) was performed as in Turner et al. (2020). Briefly, arousal state was scored using
539 a combination of cortical and hippocampal LFP, EMG, pupil diameter, and whisker and body moment. Periods of
540 manually-chosen awake rest 5s in duration with no vibrissae motion, and no body motion were used as a baseline
541 to normalize neural and hemodynamic signals. NREM sleep shows elevated cortical delta-band power and lower
542 EMG power. REM sleep is marked by elevated hippocampal theta-band power and elevated cortical gamma-band
543 power with very low baseline EMG power (muscle atonia; Cantero et al., 2004; Montgomery et al., 2008; Le Van
544 Quyen et al., 2010; Sullivan et al., 2014). Every 5 second interval was classified as either Awake, NREM sleep, or REM
545 sleep using a bootstrap aggregating random forest model with the predictors of cortical delta band power, cortical
546 beta band power, cortical gamma LFP, hippocampal theta LFP, EMG power, heart rate, and whisking duration. Sleep
547 model accuracy was validated using the out-of-bag error during model training. MATLAB functions used were
548 TreeBagger, oobError, predict.

549 **Pupil diameter and interblink interval.** Pupil diameter (**Fig. S2**) was lowpass filtered at 1 Hz with a fourth-order
550 Butterworth filter. Changes in whisking-evoked and stimulus-evoked (contralateral, auditory) diameters normalized
551 relative to the mean of the 2 second preceding the event onset. MATLAB functions used were butter, zp2sos, filtfilt.
552 Interblink interval and blink-associated physiology. Because of breaks in recording to save the data to disk, interblink
553 interval was calculated between blinks occurring within 15 min records and not blinks on the edges of trials. Blinks
554 that occurred within 1 second of each other were linked together as blinking bouts, and all blink-triggered analyses
555 were with respect to the first blink in a series. Blink-triggered averages were separated into two groups depending
556 on the arousal state classification of the 5 second bin before the blink, being either Awake (arousal state classification
557 of Awake) or Asleep (arousal state classification being either NREM or REM).

558 **Hemodynamic response function.** The hemodynamic response function (HRF, **Fig. S5**) was calculated as previously
559 described (Winder, 2017; Zhang, 2023) using both deconvolution and fitting the response to a gamma distribution.
560 Only neural activity (gamma-band power, 30-100 Hz) within 1.5 second of the stimulus was used for calculating the
561 HRF. Each HRF was calculated using half of the data and tested on the other half. To test the predictive capability of
562 each HRF, the impulse function was fit with a gamma-distribution function and convolved with the neural power to
563 predict the measured hemodynamic signal. The coefficient of determination (R^2) between the predicted and actual
564 hemodynamic signal was evaluated to quantify the efficacy of HRF's prediction (MATLAB function(s): sgolayfilt,
565 detrend, fminsearch, fitlm).

566 **Pearson's correlation coefficient calculations.** The Pearson's correlation coefficient between bilateral signals (**Fig. S6**) was obtained by mean-subtracting and digitally lowpass filtering (<1 Hz) with a fourth-order Butterworth filter
567 and then taking the Pearson's correlation coefficient (MATLAB function(s): butter, zp2sos, filtfilt, corrcoef).

569 **Arousal state transitions.** Transitions between arousal states (Awake to NREM, NREM to Awaken, NREM to REM, REM to Awake) were compared across groups (**Fig. S7**) by averaging events that met the criteria of 30 seconds of
570 one state's consecutive classifications followed by 30 seconds of another state. The difference in the $\Delta[\text{HbT}]$ between
571 the pre- and post- state transition were compared by taking the 30:10 seconds prior to the transition zero point to
572 the 10:30 seconds following. [HbT] was digitally lowpass (<1 Hz) filtered using a fourth-order Butterworth filter
573 (MATLAB function(s): butter, zp2sos, filtfilt).

575 **Data availability.** Data and code for generating the figures are available at [pending Dryad link once Manuscript
576 number is assigned] and analysis code is available at <https://github.com/KL-Turner/Turner-nNOS-Manuscript>. Raw
577 data is available upon reasonable request.

578 **Supplemental Figure 1**

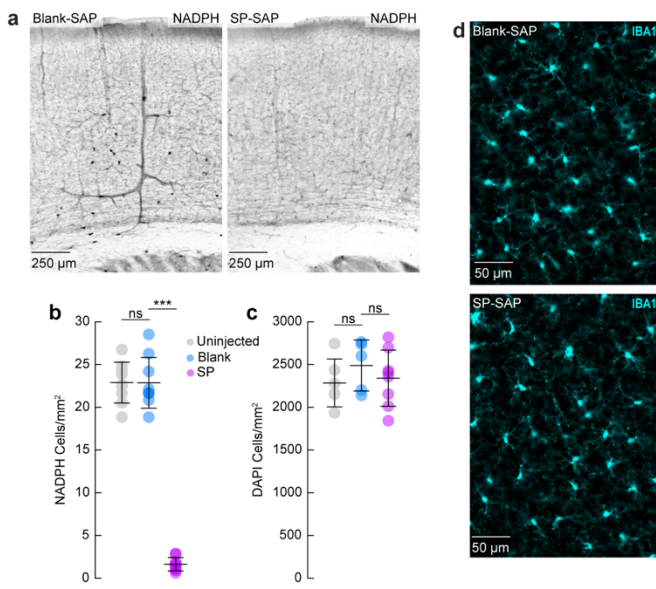


Figure S1. Histological quantification of cortical SP-Sap injections **(a)** Examples of NADPH diaphorase staining from Blank-SAP and SP-SAP injected mice. **(b)** SP-SAP injected mice (N = 9, 5M/4F) had significantly lower numbers of type-I nNOS cells than either Blank-SAP (N = 9, 4M/5F) or Uninjected mice (N = 9, 4M/5F) (SP-SAP: 1.6 ± 0.8 neurons/mm²; Blank-SAP: 22.9 ± 2.4 neurons/mm²; Uninjected: 22.9 ± 3 neurons/mm²; Blank-SAP vs. Uninjected: p = 0.98; Blank-SAP vs. SP-SAP: p = 2.23×10⁻¹³). **(c)** Counts of DAPI-labeled cell nuclei per square mm from imaged mice. Uninjected (N = 6, 6M) mice had 2284 ± 279 DAPI-labeled cell bodies/mm², which was not significantly different than those injected with Blank-SAP (N = 5, 5F, 2488 ± 298, p = 0.23, GLME), which in turn was not significantly different than those injected with SP-SAP (N = 8, 4M/4F, 2340 ± 329, p = 0.39, GLME). **(d)** Representative image of IBA1 for Blank-SAP (top) and SP-SAP (bottom) taken from **Fig 1c**. Error bars **(b, c)** denote SD. *α < 0.05, **α < 0.01, ***α < 0.001, GLME.

579 **Supplemental Figure 2**

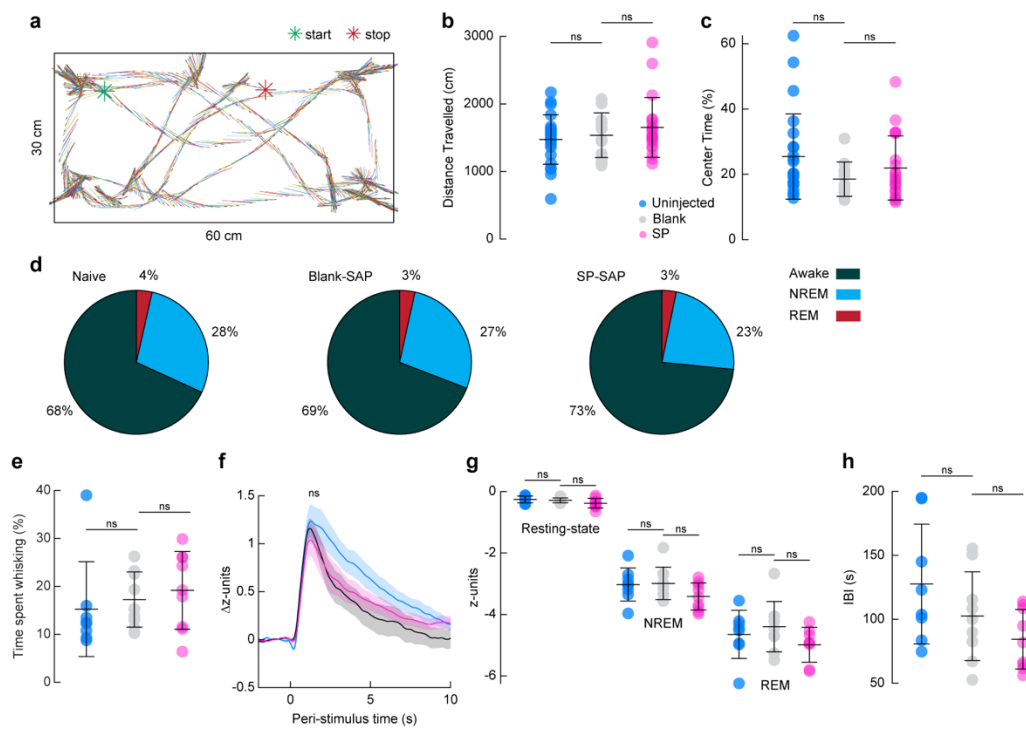


Figure S2. Behavior was unaffected by local ablation of type-I nNOS neurons **(a)** Example of exploratory behavior during 5 minutes of exploration in a novel environment. **(b)** Total distance traveled during open field exploration. Uninjected mice traveled an average of 1473.4 ± 364.7 cm compared with 1536.6 ± 329.6 cm in Blank-SAP ($p = 0.61$) and 1652.3 ± 442.7 cm in SP-SAP ($p = 0.43$). **(c)** Time spent in the center of the open field. Uninjected mice spent on average $25.5 \pm 13.1\%$ of total exploration time in the center compared with $18.5 \pm 5.3\%$ in Blank-SAP ($p = 0.08$, GLME) and $21.9 \pm 9.8\%$ in SP-SAP ($p = 0.26$). **(b, c)** $n = 12-23$ mice per group. **(d)** Percentage of time classified in each arousal state by sleep scoring. Uninjected mice spent $68.1 \pm 13.2\%$ of the time awake compared to $69.1 \pm 15.1\%$ in Blank-SAP ($p = 0.89$, ttest) and $73.4 \pm 11.6\%$ in SP-SAP ($p = 0.51$, ttest). Uninjected mice spent $28.3 \pm 11.1\%$ of the time in NREM sleep compared to $27.4 \pm 12.0\%$ in Blank-SAP ($p = 0.87$, ttest) and $23.4 \pm 9.7\%$ in SP-SAP ($p = 0.44$, ttest). Uninjected mice spent $3.5 \pm 2.9\%$ of the time in REM sleep compared to $3.4 \pm 3.5\%$ in Blank-SAP ($p = 0.96$, ttest) and $3.2 \pm 2.3\%$ in SP-SAP ($p = 0.88$, ttest). **(e)** Time spent whisking as a percentage of total imaging time. Uninjected mice spent $15.3 \pm 9.9\%$ compared with $17.3 \pm 5.8\%$ in Blank-SAP ($p = 0.61$, ttest) and $19.2 \pm 8.1\%$ in SP-SAP ($p = 0.75$, ttest). **(f)** Change in Pupil diameter in response to vibrissae stimulation. Pupil diameter of uninjected mice increased 1.3 ± 0.5 z-units following vibrissae stimulation compared with 1.2 ± 0.3 z-units in Blank-SAP ($p = 0.76$, ttest) and 1.1 ± 0.5 z-units in SP-SAP ($p = 0.50$, ttest). **(g)** Average pupil diameter across arousal states. Average pupil diameter of Uninjected mice during the resting-state was -0.25 ± 0.1 z-units compared with -0.29 ± 0.1 z-units in Blank-SAP ($p = 0.45$, ttest) and -0.34 ± 0.2 in SP-SAP ($p = 0.14$, ttest). Average pupil diameter of Uninjected mice during periods of NREM sleep was -3 ± 0.5 z-units compared with -3 ± 0.5 z-units in Blank-SAP ($p = 0.89$, ttest) and -3.4 ± 0.4 in SP-SAP ($p = 0.08$, ttest). Average pupil diameter of Uninjected mice during periods of REM sleep was -4.6 ± 0.8 z-units compared with -4.4 ± 0.8 z-units in Blank-SAP ($p = 0.53$, ttest) and -5 ± 0.6 in SP-SAP ($p = 0.11$, ttest). **(h)** Interblink-interval. Uninjected mice had a mean interblink-interval of 127.6 ± 46.8 s compared with 102.5 ± 34.7 s in Blank-SAP ($p = 0.22$, ttest) and 84.5 ± 23.3 s in SP-SAP ($p = 0.21$, ttest). **(d-h)** $n = 9$ mice per group. Shading (f) indicates SEM and error bars (b, c, e, g, h) denote SD. * $\alpha < 0.05$, ** $\alpha < 0.01$, *** $\alpha < 0.001$.

580 **Supplemental Figure 3**

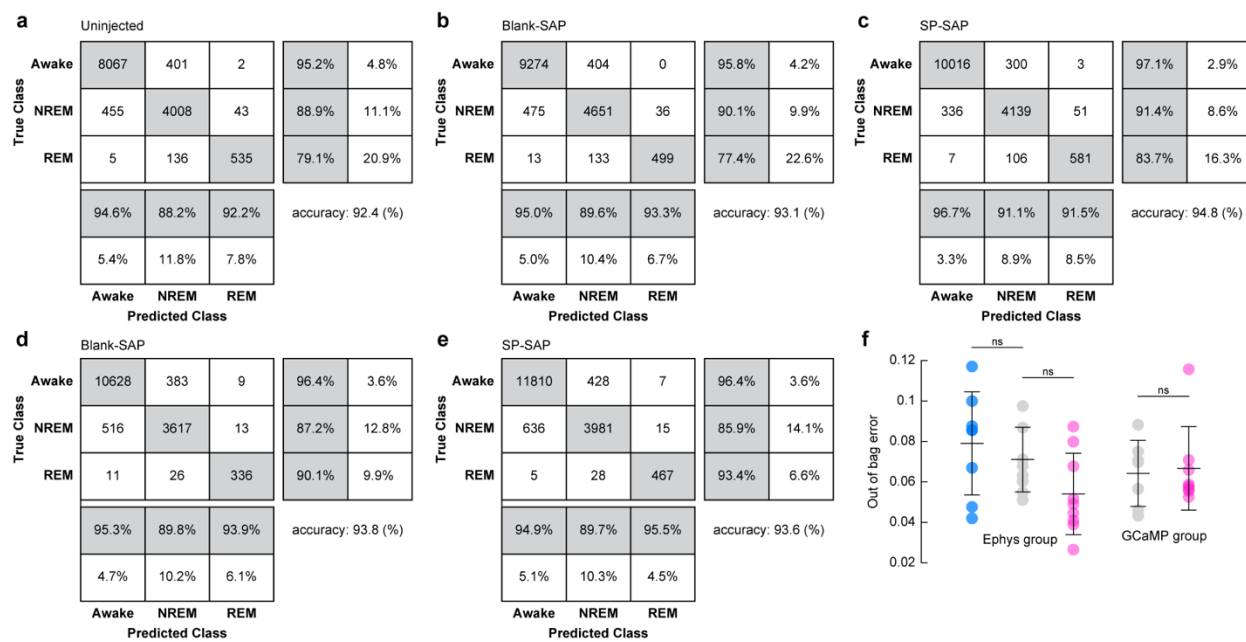


Figure S3. Sleep classification accuracy was unchanged following type-I nNOS ablation **(a)** Confusion matrix for arousal state classification of Uninjected mice with bilaterally implanted stereotrodes. **(b)** Confusion matrix for arousal state classification of Blank-SAP mice with bilaterally implanted stereotrodes. **(c)** Confusion matrix for arousal state classification of SP-SAP mice with bilaterally implanted stereotrodes. **(a-c)** n = 9 mice per group. **(d)** Confusion matrix for arousal state classification of Blank-SAP mice expressing pan-neuronal GCaMP. **(e)** Confusion matrix for arousal state classification of SP-SAP mice expressing pan-neuronal GCaMP. **(d, e)** n = 6-7 mice per group. **(f)** Out-of-bag error during training of each animal's bootstrapped random forest classification algorithm. For mice with bilateral LFP recordings, Uninjected mice had an average loss of 0.08 ± 0.03 in comparison to 0.07 ± 0.02 in Blank-SAP ($p = 0.45$, ttest) compared to 0.05 ± 0.02 in SP-SAP ($p = 0.064$, ttest). Blank-SAP mice with pan-neuronal GCaMP had an average classification loss of 0.06 ± 0.02 compared with 0.07 ± 0.02 in SP-SAP ($p = 0.80$, ttest). Error bars denote SD. * $\alpha < 0.05$, ** $\alpha < 0.01$, *** $\alpha < 0.001$.

581 **Supplemental Figure 4**

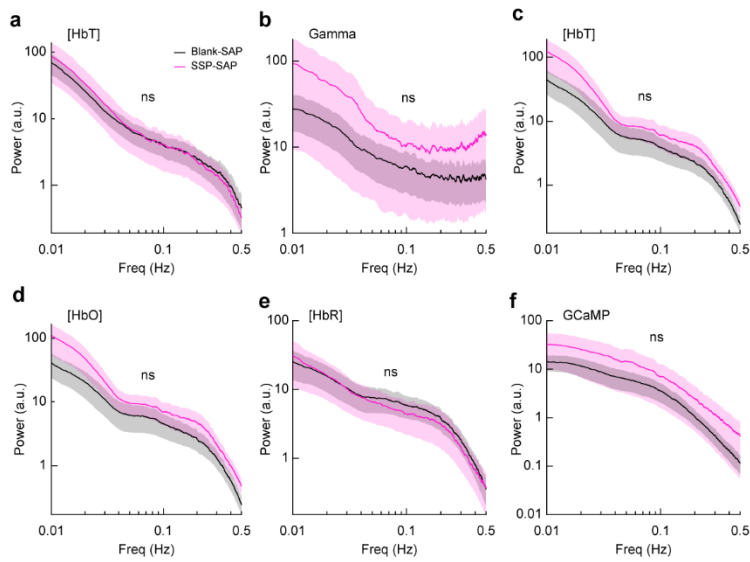


Figure S4. Ablation of Type-I nNOS neurons does not alter hemodynamic or neural power spectra Power spectral density for vascular and hemodynamic signals for the Blank-SAP group ($N = 9, 4M/5F$ for a, b; $N = 7, 3M/4F$ for c-f) and SP-SAP group ($N = 9, 5M/4F$ for a, b; $N = 8, 4M/4F$ for c-f) was not significantly different across all measurements of hemodynamic and neural signals. (a) $\Delta[HbT]$ for electrophysiology animals ($p = 0.77$, GLME). (b) Gamma-band power (second spectra) for animals with electrophysiology ($p = 0.19$, GLME). (c) $\Delta[HbT]$ for animals with GCaMP ($p = 0.27$, GLME). (d) $\Delta[HbT]$ for animals with GCaMP ($p = 0.26$, GLME). (e) $\Delta[HbT]$ for animals with GCaMP ($p = 0.68$, GLME). (f) GCaMP fluorescence ($p = 0.44$, GLME). Shading represents population averages \pm SEM. * $\alpha < 0.05$, ** $\alpha < 0.01$, *** $\alpha < 0.001$.

582 **Supplemental Figure 5**

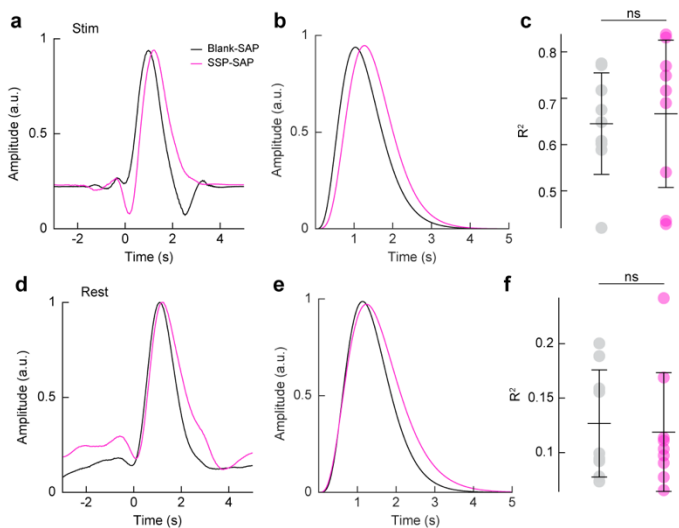


Figure S5. Ablation of Type-I nNOS neurons does not alter the predictive power of the hemodynamic response function **(a)** Stimulus-evoked hemodynamic response function obtained with deconvolution. **(b)** Stimulus-evoked hemodynamic response function fitted with a gamma distribution function. **(c)** There was no significant difference between the $\Delta[\text{HbT}]$ predictive R^2 values calculated from impulse-derived HRFs between Blank-SAP ($N = 9$, 4M/5F) 0.64 ± 0.11 or SP-SAP ($N = 9$, 5M/4F) 0.67 ± 0.16 following vibrissae stimulation ($p = 0.746$, ttest). **(d)** Resting-state hemodynamic response function based on deconvolution. **(e)** Resting-state hemodynamic response function fitted with a gamma distribution function. **(f)** There was no significant difference between the $\Delta[\text{HbT}]$ predictive R^2 values calculated from impulse-derived HRFs between Blank-SAP ($N = 9$, 4M/5F) 0.13 ± 0.05 or SP-SAP ($N = 9$, 5M/4F) 0.12 ± 0.05 during the resting-state ($p = 0.749$, ttest). Error bars denote SD. * $\alpha < 0.05$, ** $\alpha < 0.01$, *** $\alpha < 0.001$.

583 **Supplemental Figure 6**

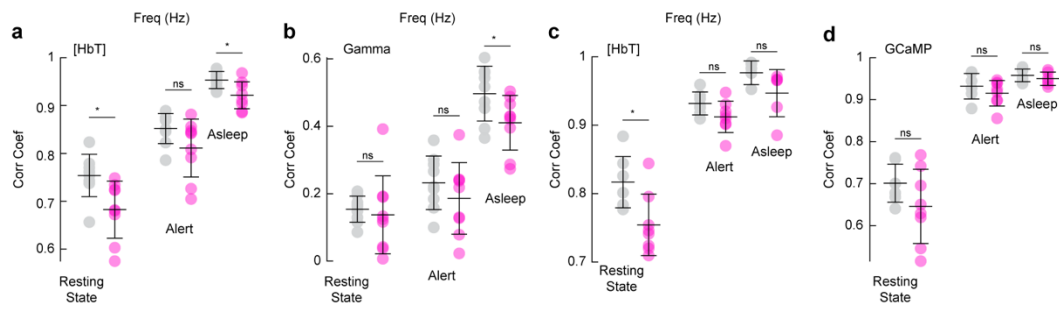


Figure S6. Pearson's correlation coefficients between bilateral hemodynamic and neural signals **(a)** Correlation coefficient between bilateral hemodynamic signals during the resting-state was 0.75 ± 0.04 in Blank-SAP and 0.68 ± 0.06 in SP-SAP (ttest $p = 0.011$), during the alert state was 0.85 ± 0.03 with Blank-SAP and 0.81 ± 0.06 in SP-SAP (ttest $p = 0.09$), and during the asleep state was 0.95 ± 0.02 in Blank-SAP and 0.92 ± 0.03 in SP-SAP (ttest $p = 0.01$). **(b)** Correlation coefficient between bilateral gamma-band power signals during the resting-state was 0.15 ± 0.04 in Blank-SAP and 0.14 ± 0.12 in SP-SAP (ttest $p = 0.69$), during the alert state was 0.23 ± 0.08 in Blank-SAP and 0.19 ± 0.11 in SP-SAP (ttest $p = 0.31$), and during the asleep state was 0.50 ± 0.08 in Blank-SAP and 0.41 ± 0.08 in SP-SAP (ttest $p = 0.04$). **(c)** Correlation coefficient between bilateral hemodynamic signals (GCaMP7s group) during the resting-state was 0.82 ± 0.04 in Blank-SAP and 0.75 ± 0.04 in SP-SAP (ttest $p = 0.01$), during the alert state was 0.93 ± 0.02 in Blank-SAP and 0.91 ± 0.02 in SP-SAP (ttest $p = 0.08$), and during the asleep state was 0.98 ± 0.02 with in-SAP and 0.95 ± 0.03 in SP-SAP (ttest $p = 0.07$). **(d)** Correlation coefficient between bilateral GCaMP7s signals during the resting-state was 0.70 ± 0.05 in Blank-SAP and 0.65 ± 0.09 in SP-SAP (ttest $p = 0.16$), during the alert state was 0.93 ± 0.03 with Blank-SAP and 0.92 ± 0.03 in SP-SAP (ttest $p = 0.31$), and during the asleep state was 0.96 ± 0.02 with Blank-SAP and 0.95 ± 0.02 in SP-SAP (ttest $p = 0.37$). Error bars denote SD. $\alpha < 0.05$, $**\alpha < 0.01$, $***\alpha < 0.001$.

584 **Supplemental Figure 7**

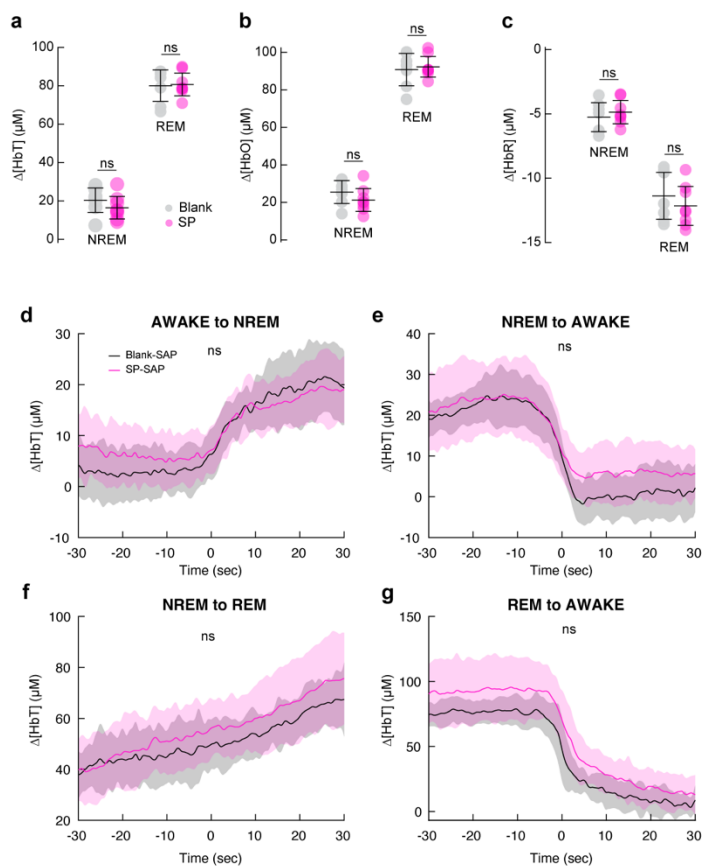


Figure S7. Removal of type-I nNOS neurons did not alter arousal state-related hemodynamic changes (a-c) Average $\Delta[\text{HbT/O/R}]$ during periods of NREM sleep and REM sleep in mice with pan-neuronal GCaMP (n = 6-7 mice per group). (a) NREM $\Delta[\text{HbT}]$ in Blank-SAP mice was 20.4 ± 6.4 μM compared to 16.5 ± 5.9 μM in SP-SAP ($p = 0.27$, ttest). REM $\Delta[\text{HbT}]$ in Blank-SAP mice was 80.0 ± 8.2 μM compared to 80.7 ± 5.9 μM in SP-SAP ($p = 0.87$, ttest). (b) NREM $\Delta[\text{HbO}]$ in Blank-SAP mice was 25.5 ± 6.1 μM compared to 21.2 ± 6.1 μM in SP-SAP ($p = 0.23$, ttest). REM $\Delta[\text{HbO}]$ in Blank-SAP mice was 90.8 ± 8.6 μM compared to 92.3 ± 5.5 μM in SP-SAP ($p = 0.72$, ttest). (c) NREM $\Delta[\text{HbR}]$ in Blank-SAP mice was -5.3 ± 1.1 μM compared to -4.9 ± 0.9 μM in SP-SAP ($p = 0.50$, ttest). REM $\Delta[\text{HbR}]$ in Blank-SAP mice was -11.4 ± 1.8 μM compared to -12.2 ± 1.5 μM in SP-SAP ($p = 0.42$, ttest). (d) Transition from Awake to NREM had a $\Delta[\text{HbT}]$ of -16.6 ± 6.4 μM in Blank-SAP mice and -11.1 ± 7.0 μM in SP-SAP mice ($p < 0.08$, GLME). (e) Transition from NREM to Awake had a $\Delta[\text{HbT}]$ of 21.5 ± 7.7 μM in Blank-SAP mice and 17.6 ± 9.5 μM in SP-SAP mice ($p < 0.33$, GLME). (f) Transition from NREM to REM had a $\Delta[\text{HbT}]$ of -17.6 ± 3.9 μM in Blank-SAP mice and -22.6 ± 7.9 μM in SP-SAP mice ($p < 0.09$, GLME). (g) Transition from REM to Awake had a $\Delta[\text{HbT}]$ of 67.8 ± 7.3 μM in Blank-SAP mice and 74.0 ± 13.1 μM in SP-SAP mice ($p < 0.21$, GLME). Error bars denote SD. Shading represents population averages \pm SEM. * $\alpha < 0.05$, ** $\alpha < 0.01$, *** $\alpha < 0.001$.

585 **Supplemental Figure 8**

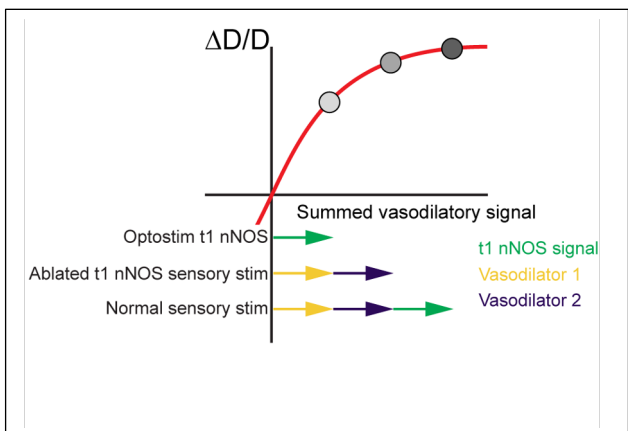


Figure S8. Schematic showing how non-linearity in the dilation response can explain coexistence strong dilation by activation of a pathway, and little change after weakening a pathway. If the diameter is a sublinear function of the sum of vasodilatory inputs, activation of all pathways will cause a dilation that is smaller than the sum of activation of each pathway individually. Loss of one pathway will not cause large changes, even though activation of that pathway in isolation can cause large dilations.

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