Thirst neurons anticipate the homeostatic consequences of eating and drinking

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Thirst motivates animals to drink in order to maintain fluid balance. Thirst has conventionally been viewed as a homeostatic response to changes in blood volume or tonicity¹⁻³. However, most drinking behaviour is regulated too rapidly to be controlled by blood composition directly, and instead seems to anticipate homeostatic imbalances before they arise⁴⁻¹¹. How this is achieved remains unknown. Here we reveal an unexpected role for the subfornical organ (SFO) in the anticipatory regulation of thirst in mice. By monitoring deep-brain calcium dynamics, we show that thirst-promoting SFO neurons respond to inputs from the oral cavity during eating and drinking and then integrate these inputs with information about the composition of the blood. This integration allows SFO neurons to predict how ongoing food and water consumption will alter fluid balance in the future and then to adjust behaviour pre-emptively. Complementary optogenetic manipulations show that this anticipatory modulation is necessary for drinking in several contexts. These findings provide a neural mechanism to explain longstanding behavioural observations, including the prevalence of drinking during meals^{10,11}, the rapid satiation of thirst^{7–9}, and the fact that oral cooling is thirst-quenching^{12–14}.

Deviations of blood volume or osmolarity from their set points are detected by specialized neurons within the circumventricular organs (CVOs) of the brain^{1–3}. Activation of these neurons generates thirst, which motivates animals to find and consume water and thereby restore fluid balance.

Nevertheless, many aspects of drinking behaviour cannot be explained by this textbook homeostatic model¹⁵ because their regulation precedes rather than responds to changes in the blood^{4–11}. For example, drinking quenches thirst tens of minutes before ingested water reaches the circulation and alters the composition of the blood, indicating that thirst is sated before homeostasis is restored. Yet animals somehow calibrate their water consumption to match their physiological need precisely^{7–9}. Conversely, most drinking occurs during meals, as a result of prandial thirst that develops long before the ingested food has been absorbed and altered the blood tonicity^{10,11}. These observations indicate that much normal drinking behaviour is anticipatory in nature, meaning that the brain predicts impending



Figure 1 | Mechanisms of homeostatic regulation of SFO^{Nos1} neurons.
a, Schematic of fibre photometry set up. Scale bar, 100 μm. b, SFO^{Nos1} neurons are activated by injection of NaCl. i.p., intraperitoneal.
c, Angiotensin blockers (INH) do not abolish the response to NaCl.
d, SFO^{Nos1} neurons are similarly activated by equiosmotic concentrations of mannitol and NaCl. e, SFO^{Nos1} neurons are activated by injection of PEG. s.c., subcutaneous. f, Angiotensin blockers abolish this response.

g, SFO^{Nos1} neurons are activated by injection of isoproterenol. **h**, Angiotensin blockers abolish this response. **i**, Schematic summarizing the mechanisms by which SFO^{Nos1} neurons monitor the blood. AngII, angiotensin-II. All error bars and shaded areas represent mean \pm s.e.m. Statistical analyses are described in Methods and Extended Data Table 1. n = 5 mice for all experiments.

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changes in fluid balance and adjusts behaviour pre-emptively. How this is achieved remains unknown.

The CVOs are the brain regions most strongly associated with fluid balance^{1–3}. Functional MRI (fMRI) studies failed to detect rapid modulation of these structures during drinking^{16,17}, leading to the idea that the integratory circuits for thirst reside in higher brain centres^{3,18} and that the CVOs function only as passive sensors of blood composition^{1–3}. However, this conclusion is complicated by the fact that the CVOs contain a diversity of intermingled neural cell types^{19,20}, which cannot be discriminated by functional imaging. Importantly, current models for the thirst circuitry have never been tested by recording the dynamics of identified neurons in awake, behaving animals.

To investigate thirst circuits *in vivo*, we focused on the SFO, a CVO that is strongly implicated in the control of drinking behaviour²¹. Optogenetic activation of SFO excitatory neurons expressing nitric oxide synthase 1 (*Nos1*) promotes voracious drinking in watersated mice^{19,22}. We replicated this (Extended Data Fig. 1) and then targeted the calcium indicator GCaMP6s to SFO^{Nos1} neurons for calcium imaging. We first confirmed in slice that GCaMP6s reported SFO^{Nos1} neuron activity induced by current injection or application of angiotensin (Extended Data Fig. 2). We then installed fibre optics above the SFO for recordings of population activity by fibre photometry²³ in awake, behaving mice (Fig. 1a).

To define the regulation of SFO^{Nos1} neurons in vivo, we recorded GCaMP fluorescence responses to a series of systemic challenges²⁴. Peripheral injection of angiotensin (Extended Data Fig. 3) or hypertonic saline (Fig. 1b) rapidly and dose-dependently activated SFO^{Nos1} neurons. The response to hypertonic saline was mediated by an osmosensory (rather than sodium-sensory) mechanism, because injection of equiosmotic concentrations of mannitol²⁵ induced a similar response (Fig. 1d). Challenge with polyethylene glycol (PEG), which induces hypovolaemia²⁶, or isoproterenol, which induces hypotension²⁴, resulted in slower and more sustained activation of SFO^{Nos1} neurons (Fig. 1e, g). The responses to PEG and isoproterenol were abolished by pre-treatment with angiotensin blockers (Fig. 1f, h), whereas the response to osmotic challenge was unaffected (Fig. 1c). Thus, SFO^{Nos1} neurons sense the osmolarity, volume and pressure of the blood via a combination of angiotensin-dependent and -independent mechanisms (Fig. 1i and Extended Data Fig. 3).

Consistent with this homeostatic regulation, overnight water restriction strongly activated SFO^{Nos1} neurons (Fig. 2a). When water was made available, mice drank avidly and, surprisingly, SFO^{Nos1} neurons were inhibited within 1 min (Fig. 2b-d). This inhibition was time-locked to the act of drinking, with activity beginning to decline the moment of the first lick (Fig. 2g) and stabilizing whenever drinking was paused. Drinking continued until the precise moment at which SFO^{Nos1} neuron activity returned to baseline, at which point drinking terminated. These responses were much too fast to be mediated by changes in the blood, which we confirmed directly by measuring plasma volume and osmolarity along a time-course of rehydration (Fig. 2e, f). Thus, contrary to current models, thirst is not quenched by the reverse of the process that generates it¹⁻³. Instead, drinking resets thirst-promoting SFO neurons in a way that anticipates the future restoration of homeostasis. Importantly, this anticipatory feedback provides a mechanism to explain how animals can match ongoing water consumption to the level of physiological need, a longstanding observation that has lacked a clear neural basis^{7–9}.

To investigate whether this rapid inhibition of SFO^{Nos1} neurons is sufficient to explain thirst satiation, we used optogenetic silencing to replay this inhibition in thirsty animals. Of note, previous studies found that SFO ablation does not completely block drinking¹, but interpretation of these lesion experiments is complicated by the fact that the SFO contains intermingled thirst-driving and -suppressing cell types^{19,20}. We targeted the inhibitory opsin eArch3.0 to SFO^{Nos1} neurons (Fig. 2h) and confirmed that this can silence SFO^{Nos1} neurons



Figure 2 | SFO Nos1 neurons receive rapid anticipatory modulation and are necessary for drinking. a, SFO^{Nos1} neurons are activated by water restriction (WR), and their activity returns to baseline (BA) after water access (WA) (n = 6 mice). **b**, Representative recording showing rapid inhibition of SFO^{Nos1} neurons during drinking after water restriction (1 out of 5 mice; red lines, licks; red boxes, bouts). c, Averaged traces showing SFO^{Nos1} neuron activity and lick rate (n = 5 mice). **d**, Quantification (n = 5 mice). **e**, Plasma osmolality (mOsm kg⁻¹) is increased by water restriction and is unchanged by 5 min after re-access (*n* = 9 [AL], 4 [-1], 9 [5], 8 [45] mice per group). AL, ad libitum. **f**, Plasma protein concentration (mg ml⁻¹) is increased by water restriction and is unchanged by 5 min after re-access (*n* = 9 [AL], 4 [-1], 9 [5], 8 [45] mice per group). g, Peristimulus time histogram (PSTH) around first lick in first bout or all other bouts (n = 5 mice). h, Schematic of optogenetic set up for silencing SFO Nos1 neurons. Scale bar, 100 $\mu m.$ i, Representative rasters of drinking after water restriction (3 out of 5 mice; black lines, licks; green boxes, laser on). j, Averaged traces showing cumulative licks following water restriction (n = 5 mice; green box, laser on). All error bars and shaded areas represent mean \pm s.e.m. Statistical analyses are described in Methods and Extended Data Table 1.

in vitro (Extended Data Fig. 4). We then water-restricted mice overnight and measured subsequent drinking behaviour. Notably, silencing SFO^{Nos1} neurons abolished drinking in dehydrated animals (Fig. 2i–j and Extended Data Fig. 4). This effect was rapidly reversible, as mice engaged in voracious drinking soon after silencing was terminated (latency to first lick 113 ± 17 s; mean \pm s.e.m.). It was also specific to drinking, as silencing did not inhibit food intake after fasting (Extended Data Fig. 7). Thus, SFO^{Nos1} neurons are necessary for water consumption, and, importantly, their rapid inhibition during drinking is sufficient to explain thirst satiation.

We sought to clarify the mechanism of rapid feedback to SFO^{Nos1} neurons during drinking. SFO^{Nos1} neuron activity was unaffected by allowing mice to see but not consume water (Fig. 3e, f) or by presenting sensory cues that had been paired with water access via Pavlovian conditioning (Extended Data Fig. 5). SFO^{Nos1} neurons are therefore not inhibited by the expectation of water availability alone, in contrast to ARC^{AgRP} neurons that control hunger^{22,27,28}. Similarly, the act of licking per se was insufficient, since SFO^{Nos1} neuron activity was not reduced in mice that performed several hundred 'air licks' after presentation of an empty water bottle (Fig. 3g, h). This indicates that a signal tightly linked to the act of water ingestion itself, such as the sensation of liquid in the oral cavity, inhibits SFO^{Nos1} neurons.

To investigate this further, we tested whether drinking would inhibit SFO^{Nos1} neurons if the ingested liquid could not restore homeostasis.



Figure 3 | Mechanisms of anticipatory regulation of SFO^{Nos1} neurons during drinking. a, Representative recording showing rapid inhibition of SFO^{Nos1} neurons during drinking after salt challenge (1 out of 8 mice; red lines, licks; red boxes, bouts). b, Averaged traces showing SFO^{Nos1} neuron activity and lick rate. c, Quantification. d, Plasma osmolality is increased by salt challenge and is unchanged by 5 min after water re-access (n = 10mice per group). e, SFO^{Nos1} neurons do not respond to the sight of water. f, Quantification. g, SFO^{Nos1} neurons do not respond to motor movements associated with licking. h, Quantification. i, SFO^{Nos1} neurons receive a

Mice were salt-challenged and then provided with access to water or hypertonic saline. Water consumption rapidly inhibited SFO^{Nos1} neurons (Fig. 3a–c and Extended Data Fig. 5), and this inhibition preceded any change in plasma osmolarity (Fig. 3d). Notably, consumption of hypertonic saline also rapidly inhibited SFO^{Nos1} neurons, with kinetics initially indistinguishable from water consumption (Fig. 3i, j). However, this initial decline was reversed after approximately 1 min. This indicates that the rapid anticipatory response to drinking has at least two components: an immediate signal that tracks fluid ingestion, and a delayed signal that reports on fluid tonicity, possibly generated by an oesophageal or gastric osmosensor.

To probe the oropharyngeal signals that regulate SFO^{Nos1} neurons further, we investigated the role of water temperature. Humans experience cold drinks as more thirst-quenching¹⁴, and a similar preference is observed in rodents^{12,13}. To test whether SFO^{Nos1} neurons contribute to this effect, we presented thirsty mice with access to water of different temperatures. We found that, regardless of temperature, mice consumed enough water to reduce the activity of their SFO^{Nos1} neurons to baseline (Fig. 3k). However, the reduction in SFO^{Nos1} neuron activity per lick was strongly temperature-dependent, with cold water inducing the largest decrease per lick and warm water the smallest (Fig. 3l). This suggests that the temperature dependence of thirst-quenching may be encoded in SFO^{Nos1} neuron activity. To test this directly, we measured the isolated effect of oral cooling on these

post-ingestive error signal that reports the osmolarity of ingested fluids. **j**, PSTH around first lick in first bout. **k**, SFO^{Nos1} neurons were similarly inhibited regardless of water temperature. **l**, Drop in activity per lick was highly temperature-dependent. **m**, Representative recording showing rapid inhibition of SFO^{Nos1} neurons during oral cooling (1 out of 5 mice; blue box, oral cooling). **n**, PSTH around placement of dry metal into oral cavity. All error bars and shaded areas represent mean \pm s.e.m. Statistical analyses are described in Methods and Extended Data Table 1. n = 5 mice for all photometry experiments.

neurons. We found that applying cold, but not room temperature, metal to the oral cavity of awake, thirsty mice was sufficient to inhibit SFO^{Nos1} neurons rapidly (Fig. 3m, n). Thus, temperature-dependent modulation of SFO^{Nos1} neurons may explain the enigmatic connection between oral cooling and thirst, including why thirsty rodents will avidly lick cold metal²⁹ and humans report that sucking on ice chips rapidly relieves thirst¹⁴.

Eating is a potent stimulus for thirst, and many animals drink primarily during meals. However, the neural basis for prandial thirst is unknown, partly because feeding stimulates drinking before any change in the blood occurs^{10,11}. To investigate the role of SFO^{Nos1} neurons, we fasted mice overnight and then provided them with access to food but not water. Remarkably, food consumption rapidly activated SFO^{Nos1} neurons, beginning at the onset of feeding and plateauing within 15 min (Fig. 4a). When water was presented, mice drank avidly and SFO^{Nos1} neurons were quickly inhibited. A similar activity pattern was observed when hungry mice were given simultaneous access to food and water (Fig. 4c). Activation of SFO^{Nos1} neurons during eating was not caused by changes in blood composition, because activation was complete before plasma osmolarity increased (Fig. 4b) and was unaffected by angiotensin blockers (Extended Data Fig. 6). Importantly, silencing SFO^{Nos1} neurons during these same protocols greatly reduced meal-associated water consumption (Fig. 4d, e) but not food intake (Extended Data Fig. 7), indicating that SFO^{Nos1} neurons are specifically necessary for prandial



Figure 4 | SFO^{Nos1} neurons are activated by eating and are required for prandial thirst. a, SFO^{Nos1} neurons are rapidly activated by eating and inhibited by prandial drinking after overnight fasting (n = 6 mice). b, Plasma osmolality is increased by eating by 45 min (but not 15 min) after chow access, and is unchanged by 5 min after water re-access (n = 10mice per group). c, SFO^{Nos1} neurons are rapidly modulated in fasted mice provided with simultaneous access to chow and water (n = 5 mice). Food refers to time (s) spent interacting with food. d, Silencing SFO^{Nos1} neurons abolishes prandial thirst in fasted mice re-fed chow before water access (n = 5 mice; green box, laser on). e, Quantification (n = 5 mice). All error bars and shaded areas represent mean \pm s.e.m. Statistical analyses are described in Methods and Extended Data Table 1.

drinking. Thus, modulation of SFO^{Nos1} neurons during feeding provides a mechanism to explain the coordination of eating and drinking, a widespread phenomenon that has lacked a clear neural substrate.

To understand better the circuit mechanisms underlying the regulation of drinking by SFO^{Nos1} neurons, we investigated the inputs and outputs of these cells. Projection mapping revealed that SFO^{Nos1} neurons densely innervate several hypothalamic nuclei associated with fluid balance³⁰ (Extended Data Fig. 8). Optogenetic stimulation of projections to the median preoptic nucleus (MnPO), but not the paraventricular hypothalamus (PVH), was sufficient to drive voracious drinking (Fig. 5a–c). Notably, examination of the monosynaptic inputs to SFO^{Nos1} neurons by retrograde rabies tracing revealed a partially overlapping set of structures (Fig. 5d–f and Extended Data Fig. 8). Determination of the cell-type-specific connectivity between these structures will be an important area for future investigation.

Ingestive behaviour is regulated by a combination of anticipatory and homeostatic cues⁴⁻⁶. Interoceptive neurons within the CVOs have been recognized for decades as crucial sites for the homeostatic regulation of drinking^{1-3,21,24,26}. By contrast, the neural substrates for the anticipatory regulation of drinking have remained obscure, despite abundant evidence that such mechanisms are important for the control of behaviour⁴⁻¹¹. Here we have demonstrated that the anticipatory signals for thirst unexpectedly converge on the same homeostatic neurons that monitor the composition of the blood (Extended Data Fig. 9). This convergence provides a straightforward mechanism for the brain to compare the needs of the body with the anticipated effects of ongoing food and water consumption and then adjust behaviour preemptively. This in turn explains longstanding behavioural observations, including the speed of thirst satiation (Fig. 2), the fact that oral cooling is thirst-quenching (Fig. 3), and the widespread coordination of eating and drinking (Fig. 4). These findings reveal that the CVOs, long viewed as merely passive sensors of the blood, receive a second class of anticipatory signals that enable their dynamic regulation of behaviour.



Figure 5 | Structure of the SFO^{Nos1} neuron-associated thirst circuit. a, Schematic of optogenetic set up for activating SFO^{Nos1} neuron somas or axon terminals in the MnPO or PVH. b, Averaged traces showing cumulative licks during photostimulation (n = 8 [SFO^{ChR2}], 6 [MnPO^{ChR2}], 7 [PVH^{ChR2}] mice per group). c, Quantification (n = 8 [SFO^{ChR2}], 5 [SFO^{GFP}], 6 [MnPO^{ChR2}], 4 [MnPO^{GFP}], 7 [PVH^{ChR2}], 4 [PVH^{GFP}] mice per group). ChR2, channelrhodopsin-2. d, Schematic of viral strategy for identifying monosynaptic inputs to SFO^{Nos1} neurons. AAV, adeno-associated virus; RV, rabies virus. e, Representative images showing SFO injection site (red, mCherry; green, green fluorescent protein; GFP) and monosynaptically connected neurons (green, GFP; blue, DAPI) in the organum vasculosum of the lamina terminalis (OVLT) and MnPO (1 out of 6 mice). Scale bars, 100 μ m. f, Quantification (n = 6 mice). ARC, arcuate nucleus; LS, lateral septum; MPA, medial preoptic area; MnR, median raphe; MS, medial septum; Pe, periventricular hypothalamus; TS, triangular septum. All error bars and shaded areas represent mean \pm s.e.m. Statistical analyses are described in Methods and Extended Data Table 1.

Online Content Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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Author Contributions C.A.Z. and Z.A.K. conceived the project and designed the experiments. C.A.Z., Y.-C.L., D.E.L., L.G., E.L.H., G.E.D. and Y.C. performed stereotaxic surgery and histology. Y.-C.L. conducted acute slice experiments. C.A.Z. and Y.C. conducted photometry experiments. C.A.Z. conducted optogenetics experiments. C.A.Z. and D.E.L. conducted plasma composition experiments. Y.C. generated the synaptophysin-GCaMP6s construct. C.A.Z., Y.-C.L., Y.C. and Z.A.K. analysed the data. C.A.Z. and Z.A.K. prepared the manuscript with input from all authors.

Author Information Reprints and permissions information is available at www.nature.com/reprints. The authors declare no competing financial interests. Readers are welcome to comment on the online version of the paper. Correspondence and requests for materials should be addressed to Z.A.K. (zachary.knight@ucsf.edu).

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METHODS

Experimental protocols were approved by the University of California, San Francisco IACUC following the NIH guidelines for the Care and Use of Laboratory Animals.

Reagents. Nos1-IRES-Cre mice (*Nos1*^{tm1(cre)}*Mgmi*], strain 017526), Agrp-IRES-Cre mice (*Agrp*^{tm1(cre)}*LowI*, strain 012899), and wild-type mice (C57BL/6J, strain 000664) were obtained from the Jackson Laboratory. Agtr1 α -GFP mice (*Tg*(*Agtr1a*-*EGFP*)*NZ44Gsat*, strain 033059) were obtained from MMRRC. Adult mice (>4 weeks old) of both genders were used for experiments.

Recombinant AAVs expressing ChETA_{TC} (AAV5-EF1 α -DIO-hChR2(E123T/T159C)-P2A-mCherry; AAV5-CaMK2 α -hChR2(E123T/T159C)-P2A-mCherry), ChR2/H134R (AAV5-CaMK2 α -hChR2(H134R)-YFP), eArch3.0 (AAV5-EF1 α -DIO-eArch3.0-YFP), mCherry (AAV5-EF1 α -DIO-mCherry), and GFP (AAV5-CaMK2 α -GFP) were obtained from the UNC Vector Core. Recombinant AAVs expressing GCaMP6s (AAV1-hSyn-FLEX-GCaMP6s; AAV5-hSyn-FLEX-GCaMP6s) and eArch3.0 (AAV5-CBA-FLEX-eArch3.0-GFP) were obtained from the Penn Vector Core. Recombinant EnvA-pseudotyped G-deficient rabies virus expressing GFP (RV-EnvA- Δ G-GFP) was obtained from the Salk Institute.

Plasmids encoding synaptophysin-GCaMP6s and GFP-RPL10a fusion proteins were generated in-house. Recombinant AAVs containing these plasmids (AAV5-EF1 α -DIO-synaptophysin-GCaMP6s; AAV2-EF1 α -FLEX-GFP-RPL10a) were commercially produced by the UNC Vector Core. Rabies helper viruses (AAV1-CAG-DIO-TVA-mCherry; AAV1-CAG-DIO-G) were obtained from the laboratory of N. Shah. **Stereotaxic surgery.** For SFO injections, 100–300 nl of virus was injected at one site (-0.50 mm antero-posterior (AP); 0 mm medio-lateral (ML); -2.75 mm dorso-ventral (DV) relative to bregma). For ARC injections, 1 μ l of virus total was injected at two sites (-1.85 mm AP; -0.3 mm ML; -5.7 and -5.8 mm DV). Mice were allowed 2–4 weeks for viral expression and recovery from surgery before behavioural testing.

For soma photostimulation experiments, recombinant AAVs expressing ChETA_{TC}, mCherry, or GFP were stereotaxically injected into the SFO of Nos1-IRES-Cre mice and wild-type mice. In the same surgery, a fibreoptic cannula was implanted unilaterally above the SFO (-2.25 mm DV).

For terminal photostimulation experiments, recombinant AAVs expressing hChR2/H134R or GFP were injected into the SFO of wild-type mice. In the same surgery, a fibreoptic cannula was implanted unilaterally above the MnPO (+0.45 mm AP; +0 mm ML; -3.7 mm DV) or PVH (-0.75 mm AP; +0.2 mm ML; -3.9 mm DV).

For photoinhibition experiments, recombinant AAVs expressing eArch3.0 or mCherry were injected into the SFO of Nos1-IRES-Cre mice. In the same surgery, a fibreoptic cannula was implanted unilaterally above the SFO (-2.35 mm DV).

For photometry experiments, recombinant AAVs expressing GCaMP6s or GFP were injected into the SFO of Nos1-IRES-Cre mice and into the ARC of Agrp-IRES-Cre mice. In the same surgery, a fibreoptic cannula was implanted unilaterally above the SFO (-2.95 mm DV) or ARC (-5.7 mm DV).

For projection-mapping experiments, recombinant AAVs expressing synaptophysin-GCaMP6s were injected into the SFO of Nos1-IRES-Cre mice. After four weeks, mice were euthanized and processed for histology.

For rabies tracing experiments, recombinant AAVs expressing TVA and G were injected into the SFO of Nos1-IRES-Cre mice. After two weeks, recombinant EnvA-pseudotyped G-deficient rabies virus expressing GFP was injected into the SFO. After one additional week, mice were euthanized and processed for histology.

For retrobeads tracing experiments, red RetroBeads IX (LumaFluor) were injected into the SFO of wild-type mice. After one week, mice were euthanized and processed for histology.

Slice electrophysiology and calcium imaging. Acute forebrain slices were prepared from 8-15-week-old Nos1-IRES-Cre mice expressing ChETA_{TC}, eArch3.0, or GCaMP6s for 2-4 weeks. Fluorescent cells in the SFO were identified for whole-cell patch clamp recordings. Slices were sectioned in ice-cold oxygenated (95% O₂/5% CO₂) cutting saline containing (in mM) 26 NaHCO₃, 1.25 NaH₂PO₄, 3 KCl, 10 glucose, 210 sucrose, 2 CaCl₂, 2 MgCl₂. Slices were then transferred to oxygenated artificial cerebrospinal fluid (ACSF) containing (in mM) 125 NaCl, 25 NaHCO₃, 1.25 NaH₂PO₄, 2.5 KCl, 15 glucose, 2 CaCl₂, 1 MgCl₂ and incubated at 34 °C for 30 min. Slices were then stored at room temperature until used. During experiments, slices were placed in a recording chamber and superfused with oxygenated ACSF. Glass pipettes for recording $(3-8 M\Omega)$ were pulled from borosilicate glass capillary (O.D. 1.5 mm, I.D. 0.86 mm, Sutter Instrument) and filled with internal solution containing (in mM) 125 K gluconate, 10 KCl, 4 NaCl, 4 Mg₃ATP₂, 0.3 Na₃ATP, 5 Na₂-phosphocreatine, 10 HEPES. Whole-cell recordings were made at 28 °C using an Axopatch 700B amplifier (Molecular Devices). Data acquisition (filtered at 5 kHz and digitized at 10 kHz) and pulse generation were performed using a Digidata 1550 (Molecular Devices) and pClamp software (version 10.5, http://www.moleculardevices.com).

For channelrhodopsin validation, cells were photostimulated under current clamp mode using an LED light source (Lambda HPX, Sutter Instruments) pulsing light at 5–40 Hz through a 470 nm excitation filter set (U-N41 017, E.X. 470 nm, B.S. 495 nm, E.M. 5, Olympus).

For archaerhodopsin validation, cells were activated using currents ranging from 0 pA to 45 pA (ΔI = 5 pA) for 3 s in duration injected under current clamp mode. During current injection, cells were photosilenced using an LED light source (Lambda HPX, Sutter Instruments) sending constant light through a 500 nm excitation filter set (49003, E.X. 500 nm, B.S. 515 nm, E.M. 5, Chroma Technology).

For calcium imaging validation, cells were activated using currents (30 pA) ranging from 100 ms to 1 s (Δt = 100 ms) in duration injected under current clamp mode. Calcium imaging was performed simultaneously using a digital CCD camera (Hamamatsu, ORCA-ER) mounted on an upright microscope (Olympus, BX51). Micro-manager software (version 1.4, https://www.micro-manager.org) was used as microscope control interface. After loading, cells were imaged (10 ms exposure time; 10 Hz) using an LED light source (Lambda HPX, Sutter Instruments) sending constant light through a 470 nm excitation filter set (U-N41 017, E.X. 470 nm, B.S. 495 nm, E.M. 5, Olympus). We have previously validated the use of GCaMP6s to image calcium signals in ARC^{AgRP} neurons²⁷.

To visualize the responsiveness of SFO^{Nos1} neurons to angiotensin-II (AngII), 1 µM AngII was bath-applied using a slow perfusion system (2 ml min⁻¹) during imaging. Data analysis followed the basic logic described previously². Regions of interest (ROIs) were selected using the polygon selection tool in ImageJ with cell nucleus included. The plot *z*-axis profile function in ImageJ was then used to measure the mean value of each ROI versus frame number. Neuropil fluorescence was selected and estimated using the same protocol. Only regions located near the cell with no detectable fluorescent neural processes were used. The true fluorescence signal of each cell was estimated with function²⁷: $F_{cell_true}(t) = F_{cell_measured}(t) - r \times F_{neuropil}(t)$, where r=0.7.

Photostimulation *in vivo.* All experiments were performed in behavioural chambers (Coulbourn Instruments, Habitest Modular System). Experiments were performed during the light cycle to control for circadian factors and performed in a well-lit environment illuminated with white light. Water consumption was monitored with an optical lickometer (Colbourn Instruments). Mice were acclimated to the behavioural chamber for at least 15 min before the beginning of each testing session. For some experiments, video was recorded using cameras installed above each cage.

To test whether acute photostimulation could induce drinking, mice were provided with constant access to water and monitored for 30 min (pre-stim), then photostimulated for 30 min (stim, 10 ms pulses at 20 Hz for 1 s every 4 s, 20–25 mW) using a DPSS 473 nm laser (Shanghai Laser and Optics Century), and then monitored for another 30 min (post-stim).

For saline drinking experiments, the same model was used except that mice were provided with access to 150 mM NaCl in water.

For delayed-access experiments, the same model was used except that stimulation lasted 60 min instead of 30 min. Water access was removed after 30 min pre-stimulation period and water re-access was provided 30 min later.

For water restriction experiments, mice were water-restricted for 24 h in their home cages, acclimated to the behavioural chamber for 15 min, and then provided with access to water for 30 min. To compare photostimulation to water restriction, all sessions were aligned to the first lick in the stimulation or water-access period.

For food consumption experiments, mice were fed ad libitum in their home cages, acclimated to the behavioural chamber for 15 min, and then provided with a single pellet of chow without simultaneous access to water for 30 min. After 30 min, chow was removed and the amount consumed measured.

Photoinhibition *in vivo*. All experiments were performed in behavioural chambers (Coulbourn Instruments, Habitest Modular System). Experiments were performed during the light cycle to control for circadian factors and performed in a well-lit environment illuminated with white light. Water consumption was monitored with an optical lickometer (Colbourn Instruments). Mice were acclimated to the behavioural chamber for at least 15 min before the beginning of each testing session.

To test whether acute silencing could inhibit drinking, mice were waterrestricted for 24 h in their home cages, acclimated to the behavioural chamber for 15 min, and then provided with access to water for 15 min with or without optical silencing (10–15 mW) using a DPSS 532 nm laser (Shanghai Laser and Optics Century), then monitored for another 15 min without optical silencing. Trials with and without photoinhibition ('–laser' and '+laser') were interleaved.

To test whether acute optical silencing could inhibit prandial drinking in fasted mice, mice were fasted for 24h in their home cages, acclimated to the behavioural chamber for 15 min, then provided with a single pellet of chow under one of two experimental conditions: (1) no water access and no laser for 45 min, chow removal, then water access \pm laser for 30 min, (2) no water access \pm laser for 2h. In experimental condition 2, chow was removed and the amount consumed measured every 15 min.

Fibre photometry. Construction of the rig for performing fibre photometry has been previously described²⁷. The signal was output to a lock-in amplifier (Stanford Research System, SR810) with time constant 30 ms to allow filtering of noise at higher frequency. Signal was then digitized with LabJack U6-Pro and recorded using software provided by LabJack (http://labjack.com/support/software) with 250 Hz sampling rate.

All experiments were performed in behavioural chambers (Coulbourn Instruments, Habitest Modular System). Experiments were performed during the light cycle to control for circadian factors and performed in a well-lit environment illuminated with white light. Mice were acclimated to the behavioural chamber for at least 15 min with access to 24 °C water before the beginning of each testing session. Photometry data were subjected to minimal processing consisting of only within-trial fluorescence normalization. For some experiments, video was recorded using cameras installed above each cage.

For pharmacological experiments, mice were acclimated to the behavioural chamber for 15 min, then given an injection and monitored for 45 min. AngII ($20 \mu g$ per mouse, $200 \mu g$ per mouse), losartan ($100 \, mg \, kg^{-1}$), captopril ($50 \, mg \, kg^{-1}$), and PEG (40%) were administered subcutaneously, and NaCl ($1 \, M, 2 \, M, 3 \, M$), mannitol ($2 \, M$), and isoproterenol ($100 \, mg \, kg^{-1}$) were administered intraperitoneally. All subcutaneous injections were given in a total volume of $400 \, \mu$ l with PBS as vehicle, and all intraperitoneal injections were given in a total volume of $150 \, \mu$ l with PBS as vehicle. To block angiotensin signalling, losartan was administered 30 min before AngII injection, and losartan plus captopril (described as 'angiotensin blockers' and 'INH') were administered simultaneously to PEG/NaCl/isoproterenol injection or chow access. Losartan is a selective AngII type 1 receptor (AT_1R) antagonist³¹, and captopril is an angiotensin converting enzyme (ACE) inhibitor³². Mice were not provided with access to water unless otherwise noted.

For water restriction experiment in Fig. 2a, mice were placed in the behavioural chamber and calcium signals recorded for 10 min on day one ('BA'). Mice were then returned to their home cages and water-restricted for 24 h. After 24 h of water restriction, mice were placed in the behavioural chamber and calcium signals again recorded for 10 min on day two ('WR'). Mice were then returned to their home cages and immediately provided with access to water. After 1 h of water re-access, mice were placed in the behavioural chamber and calcium signals again recorded for 10 min ('WA'). Photometry settings, including laser power and time constant, were the same for every mouse and every recording session. The reported fluorescence was calculated as the median fluorescence of minutes 5–10 of each recording.

For other water restriction experiments, mice were water-restricted for 48 h in their home cages, acclimated to the behavioural chamber for 15 min, and then provided with access to water for 45 min. For all experiments, the opening of a guillotine port cued water access.

For salt-loading experiments, mice were acclimated to the behavioural chamber for 15 min, and then given an intraperitoneal injection of 150 μ l 3 M NaCl. After 45 min, mice were provided with access for 45 min to a bottle that was empty, contained 12 °C, 24 °C or 36 °C water, or contained 24 °C 300 mM NaCl in water. Quantification of PSTHs in Fig. 3f, h refers to $\Delta F/F$ at 15 s. The PSTHs in Fig. 3g represent the first bout from 24 °C water bottle and all bouts from empty bottle.

For oral cooling experiments, mice were given an intraperitoneal injection of $150\,\mu$ l 3 M NaCl. After 10–15 min, a piece of dry metal (ice-cold, 'oral cooling'; room temperature, 'sham') was placed in the oral cavity, held for 30 s, and then removed. This process was repeated after >60 s wait with metal of the other temperature. The temperature order was counter-balanced across trials.

For sucrose drinking experiments, mice were provided with access to food and water ad libitum before testing. Mice were acclimated to the behavioural chamber for 15 min, then provided with ad libitum access to 150 mM sucrose for >2 h.

For fasting–refeeding experiments, mice were fasted for 24 h in their home cages, acclimated to the behavioural chamber for 15 min, and then provided with a single pellet of chow either with or without simultaneous access to water for 45 min. After 45 min, chow was removed and the amount consumed measured, and mice were immediately provided with access to water for 45 min.

For Pavlovian conditioning experiments, mice were acclimated to the behavioural chamber for 15 min, and then given an intraperitoneal injection of 150 μ l 3 M NaCl. After 45 min, an auditory cue was played (2.9 kHz, 300 ms; Colbourn Instruments), and 3 s later the water port was opened and mice were provided with access to water for 45 min. There were seven training trials and a final probe trial (both training and probe trials included auditory cue).

Data analysis. All data were analysed with custom Matlab code. Throughout the paper, a drinking bout is defined as any set of ten or more licks in which no interlick interval is greater than one second.

For photometry data, all responses were normalized to baseline using the function: $\Delta F/F = (F - F_0)/F_0$, in which F_0 is the median fluorescence of the baseline period. The baseline period for full experiments was 15 min before time zero, and the baseline period for PSTHs around drinking bouts was 60 s before

the first lick or 60 s after the last lick in a bout (15 s for cue and empty bottle plots). Time zero was defined as the moment the mouse was returned to the behavioural chamber following injection, the moment of water access, or the moment of chow access. The time constant, τ , was estimated as described previously²⁷. In bar graphs quantifying $\Delta F/F$, the median fluorescence of a 1-s window around the indicated time is reported, and 1 min before time zero is reported as baseline.

For rabies tracing experiments, mice were transcardially perfused with PBS followed by formalin. Brains were post-fixed overnight in formalin and placed in 20% sucrose for 24h. Free-floating sections ($40\,\mu m$) were prepared with a cryostat and half of sections were immediately mounted and imaged by direct fluorescence with a Zeiss LSM 700 confocal microscope. Quantification was performed using the cell counter tool in ImageJ.

Plasma osmolality. For water restriction experiments, wild-type mice were water-restricted for 24 h, then provided with access to water at t = 0. For salt-loading experiments, wild-type mice were given an intraperitoneal injection of 150µl 3 M NaCl at t = 0, and then provided with access to water at t = 45 min. For fasting-refeding experiments, wild type mice were fasted for 24 h, then provided with a single pellet of chow without simultaneous access to water at t = 0. At t = 45 min, chow was removed, and mice were immediately provided with access to water. At a single time point per session (hydrated, 0 min, 5 min, 45 min for water restriction experiments; 0 min, 5 min, 50 min for fasting-refeding experiments), 125µl of blood was collected from the tail vein using EDTA-coated capillary tubes (RAM Scientific). Plasma was isolated by centrifugation, and osmolality was measured using a freezing point osmometer (Fiske Associates). Mice were allowed 1 week for recovery between sessions.

Plasma protein. Wild-type mice were water-restricted for 24 h, and then provided with access to water at t = 0. At a single time point per session (hydrated, 0 min, 5 min, 45 min), 125 µl of blood was collected from the tail vein using EDTA-coated capillary tubes (RAM Scientific). Plasma was isolated by centrifugation, and plasma protein concentration was measured using a BCA protein assay kit (Thermo Fisher Scientific). Mice were allowed 1 week for recovery between sessions. Plasma protein concentration is inversely proportional to plasma volume.

Immunohistochemistry. Mice were transcardially perfused with PBS followed by formalin. Brains were post-fixed overnight in formalin and placed in 20% sucrose for 24 h. Free-floating sections (40μ m) were prepared with a cryostat, blocked (3% BSA, 2% NGS, and 0.1% Triton-X in PBS for 2 h), and then incubated with primary antibody (chicken anti-GFP, Abcam, ab13970, 1:1,000; rat anti-RFP, ChromoTek, 5f8, 1:2,000; rabbit anti-cFos, Santa Cruz Biotech, sc52, 1:1,000) overnight at 4 °C (two nights for cFos staining). Sections were then washed, incubated with secondary antibody (Alexa Fluor 488 goat anti-chicken, Life Technologies, a11039, 1:1,000; Alexa Fluor 568 goat anti-rabbit, Life Technologies, a11011, 1:1,000) for 2 h at room temperature, mounted, and imaged with a Zeiss LSM 700 confocal microscope. Sections stained for cFos underwent unmasking before blocking (1% H₂O₂ and 1% NaOH in PBS for 10 min; 0.3% glycine in PBS for 10 min; 0.03% SDS in PBS for 10 min).

Statistics. Values are reported as mean \pm s.e.m. (error bars or shaded area). The shaded area in Extended Data Fig. 2d represents the 95% confidence interval for the line-of-best-fit. Statistical analyses and linear regressions were performed using Matlab or Prism. P values for pair-wise comparisons were performed using a two-tailed Student's t-test. P values for comparisons across multiple groups were performed using analysis of variance (ANOVA) and corrected for multiple comparisons using the Holm–Šídák method. *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001. Result sheets of statistical tests from Prism detailing (wherever applicable) estimates of variance within each group, comparison of variances across groups and so on are available on request. Animals for fibre photometry experiments were excluded if there was no response (<10%) to 3 M NaCl injection i.p. (SFO^{Nos1} neurons) or to 60 µg ghrelin injection i.p. (ARC^{AgRP} neurons). Animals for optogenetics experiments were excluded based on histology (expression of ChETA_{TC}/hChR2(H134R)/eArch3.0 in SFO) and fibreoptic placement. We observed few and sparse virally infected cells outside the SFO. These criteria were pre-established. No statistical method was used to predetermine sample size. Randomization and blinding were not used.

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Extended Data Figure 1 | See next page for caption.



Extended Data Figure 1 | Optogenetic activation of SFO^{Nos1} neurons is sufficient to promote drinking, but negative feedback inhibits excessive drinking during optogenetically and dehydration-induced drinking. a-l, Optogenetic activation of SFO^{Nos1} neurons rapidly and specifically promotes drinking. a, Expression of mCherry in SFO^{Nos1} neurons from AAV5-EF1α-DIO-ChETA_{TC}-2A-mCherry. Scale bar, 100 μm. **b**, Representative recording showing rapid firing of SFO^{Nos1} neuron in response to photostimulation (20 Hz) in acute slice (1 out of 3 cells; blue lines, stimulation). c, Schematic of optogenetic set up for activating SFO^{Nos1} neurons. **d**, Rasters of drinking in response to optogenetic stimulation for seven trials each for four SFO^{Nos1}::ChETA_{TC} mice (black lines, licks; blue boxes, stimulation). e, Averaged traces showing lick rate $(n = 6 \text{ SFO}^{\text{Nos1}}$::mCherry mice and 8 SFO^{Nos1}::ChETA_{TC} mice). **f**, Averaged traces showing cumulative licks (n = 6 SFO^{Nos1}::mCherry mice and 8 SFO^{Nos1}::ChETA_{TC} mice). g, Quantification of drinking during stimulation protocol (****P < 0.0001, two-way repeated-measures ANOVA, n = 6 SFO^{Nos1}::mCherry mice and 8 SFO^{Nos1}::ChETA_{TC} mice). h, Licks during stimulation period across seven consecutive trials (not significant (n.s.), one-way repeated-measures ANOVA, n = 8 mice). i, Latency to first lick during stimulation period across seven consecutive trials (n.s., one-way repeated-measures ANOVA, n = 8 mice). j, Heat maps showing location of SFO^{Nos1}::ChETA_{TC} mice during stimulation protocol (n = 4 mice). k, Quantification of time spent at lickometer during stimulation protocol (**P < 0.01, one-way repeated-measures ANOVA, n = 4 mice). I, Activation of SFO^{Nos1} neurons did not induce feeding (n.s., two-way repeated-measures ANOVA, n = 6 SFO^{Nos1}::mCherry mice and 4 SFO^{Nos1}::ChETA_{TC} mice). Panels m-q show that osmotic dilution does not inhibit excessive drinking during optogenetically induced

drinking. m, SFO^{Nos1}::ChETA_{TC} mice were stimulated, but with access to 150 mM NaCl instead of water. **n**, Lick rate (n = 8 mice). **o**, Cumulative licks (n = 8 mice). **p**, Comparison to stimulation with water access (n = 8mice). **q**, Quantification (n.s., two-tailed Student's *t*-test, n = 8 mice). Panels **r**-**v** show that channelrhodopsin failure does not explain the negative feedback that inhibits excessive drinking during optogenetically induced drinking. r, SFO^{Nos1}::ChETATC mice were stimulated, but with delayed access to water instead of immediate access. s, Lick rate (n = 4mice). t. Cumulative licks (n = 4 mice). u. Comparison to stimulation with water access (n = 4 mice). v, Quantification (n.s., two-tailed Student's *t*-test, n = 4 mice). Panels w-af show that a negative feedback mechanism also inhibits excessive drinking during dehydration-induced drinking. w, Comparison of optogenetically and dehydration-induced drinking in SFO^{Nos1}::ChETA_{TC} mice (n = 4 mice; P value colour bar represents independent two-tailed Student's t-tests). x, Latency to first lick (n.s., twotailed Student's *t*-test, n = 4 mice). y, Cumulative licks (*P < 0.05, twoway repeated-measures ANOVA, n = 4 mice). z, Cumulative probability distribution for inter-lick interval, a measure of licking 'speed' (n = 4mice). aa, Median inter-lick interval (**P < 0.01, two-tailed Student's *t*-test, n = 4 mice). **ab**, Time constant (τ) for cumulative licks (*P < 0.05, two-tailed Student's *t*-test, n = 4 mice). ac, Number of drinking bouts (**P < 0.01, two-way repeated-measures ANOVA, n = 4 mice). ad, Number of licks per drinking bout (**P < 0.01, two-way repeatedmeasures ANOVA, n = 4 mice). **ae**, Bout duration (**P < 0.01, two-way repeated-measures ANOVA, n = 4 mice). af, Inter-bout interval (n.s., twoway repeated-measures ANOVA, n = 4 mice). Values are mean \pm s.e.m. (error bars or shaded area).



Extended Data Figure 2 | GCaMP6s accurately reports SFO^{Nos1} neuron activity in acute slices. a, Expression of GCaMP6s in SFO^{Nos1} neurons from AAV5-EF1 α -FLEX-GCaMP6s. Scale bar, 100 µm. b, Representative fluorescence images of a neuron given a 30 pA current injection for 700 m in acute slice (1 out of 9 cells). c, Representative traces showing calcium responses in response to 30 pA current injections of increasing duration to produce increasing numbers of action potentials (1 out of 9 cells). d, Relationship between number of action potentials and $\Delta F/F$ for the representative neuron in c (shaded area denotes 95% confidence interval). e, R^2 and P value for linear relationship between number of action potentials and $\Delta F/F$ (n = 9 cells). Panels f–j show that SFO^{Nos1} neurons are homogeneously responsive to both angiotensin and salt challenge. f, Representative fluorescence images showing SFO^{Nos1} neuron activity before and during bath application of angiotensin (1 out of 3 experiments;

red circles, identified neurons). **g**, 24 out of 27 (~90%) identified SFO^{Nos1} neurons were activated by bath application of angiotensin (red line, mean; grey lines, individual activated neurons). **h**, Quantification (*****P* < 0.0001, two-way repeated-measures ANOVA, *n* = 24 activated neurons). **i**, Experimental design to test whether a single population of SFO neurons is responsive to both angiotensin and salt challenge. **j**, Co-localization of Agtr1 α ::GFP and salt-challenge-induced cFos indicates that SFO^{Nos1} neurons are homogeneously responsive to both angiotensin and salt challenge. Scale bars, 100 µm. **k**, Experimental design to test whether a SFO neuron marker CaMK2 α . **l**, Co-localization of CaMK2 α ::mCherry and Nos1::GFP indicates that SFO^{Nos1} neurons are excitatory. Scale bar, 100 µm. Values are mean \pm s.e.m. (error bars or shaded area).





Extended Data Figure 3 | **Regulation of SFO**^{Nos1} **neurons by homeostatic signals. a**, Recordings from SFO^{Nos1}::GCaMP6s mice as they explored a behavioural chamber without access to food or water revealed dynamic fluctuations in fluorescence around a stable baseline (1 out of 8 mice); these fluctuations were absent from recordings from SFO^{Nos1}::GFP mice (1 out of 3 mice). **b**, Quantification of response to peripheral injection of NaCl (averaged traces in Fig. 1b; **P < 0.01, ****P < 0.0001, two-way repeated-measures ANOVA, n = 5 mice). **c**, Time constant (τ) of rising and falling phases of response to peripheral injection of NaCl (n.s., two-way repeated-measures ANOVA, n = 5 mice). **d**, Representative recordings for five mice showing response to peripheral injection of NaCl or vehicle. **e**, SFO^{Nos1} neurons are activated by peripheral injection of angiotensin in a dose-dependent manner (n = 3 mice). **f**, Quantification (*P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001, two-way repeatedmeasures ANOVA, n = 3 mice). **g**, The AT₁R antagonist losartan abolished

the response to peripheral injection of angiotensin (n = 3 mice). **h**, Quantification (**P < 0.01, two-way repeated-measures ANOVA, n = 3mice). **i**, Schematic illustrating expected observations whether activation of SFO^{Nos1} neurons in response to peripheral injection PEG/isoproterenol and NaCl is angiotensin-dependent. **j**, Angiotensin blockers abolished the response to peripheral injection of PEG (quantification in Fig. 1f; n = 5 mice). **k**, Angiotensin blockers abolished the response to peripheral injection of isoproterenol (quantification in Fig. 1h; n = 5 mice). **l**, Angiotensin blockers did not abolish the response to peripheral injection of NaCl (quantification in Fig. 1c; n = 5 mice). **m**, Schematic illustrating expected observations if activation of SFO^{Nos1} neurons in response to peripheral injection NaCl is sodium-sensory or osmosensory. **n**, SFO^{Nos1} neurons are similarly activated by peripheral injection of equimolar concentrations of mannitol and NaCl (quantification in Fig. 1d; n = 5mice). Values are mean \pm s.e.m. (error bars or shaded area).





Extended Data Figure 4 | **SFO**^{Nos1} **neurons are necessary for drinking. a**, Expression of yellow fluorescent protein (YFP) in SFO^{Nos1} neurons from AAV5-EF1 α -DIO-eArch3.0-YFP. Scale bar, 100 μ m. **b**, Representative recording showing firing of SFO^{Nos1} neurons is blocked in response to photosilencing in acute slice (1 out of 3 cells; yellow line, laser). **c**, Averaged traces showing lick rate for experiment in Fig. 2j (n = 5 mice; green box, laser on). **d**, Quantification (*P < 0.05, two-way repeated-

measures ANOVA, n = 5 mice). **e**, Averaged traces showing cumulative licks after water restriction for SFO^{Nos1}::mCherry control mice (n = 5 mice; green box, laser on). **f**, Quantification (n.s., two-way repeated-measures ANOVA, n = 5 mice). **g**, Averaged traces showing lick rate after water restriction for SFO^{Nos1}::mCherry control mice (n = 5 mice; green box, laser on). **h**, Quantification (n.s., two-way repeated-measures ANOVA, n = 5 mice). Values are mean \pm s.e.m. (error bars or shaded area).

RESEARCH LETTER



Extended Data Figure 5 | Regulation of SFO^{Nos1} neurons by anticipatory signals. a, Representative recordings for five mice showing activation of SFO^{Nos1} neurons during salt challenge and rapid inhibition of SFO^{Nos1} neurons during drinking. **b**, PSTH of SFO^{Nos1} neuron activity and lick rate around the first lick in either the first drinking bout or all other drinking bouts following salt challenge (n = 5 mice). c, The decrease in SFO^{Nos1} neuron activity was greatest during the first drinking bout ($\Delta F/F$ at 20 s after first lick; **P < 0.01, two-tailed Student's *t*-test, n = 5 mice). **d**, PSTH of SFO^{Nos1} neuron activity and lick rate around the last lick in either the first drinking bout or all other drinking bouts following salt challenge (n = 5 mice). **e**, Representative recording showing no inhibition of SFO^{Nos1} neurons during licking an empty bottle following salt challenge (1 out of 5 mice; red lines, licks; red boxes, drinking bouts). f, Representative recording showing rapid inhibition followed by 're-setting' of SFO^{Nos1} neurons during drinking 300 mM NaCl following salt challenge (1 out of 5 mice; red lines, licks; red boxes, drinking bouts). g, SFO^{Nos1} neurons receive a post-ingestive error signal that reports the osmolarity of ingested

fluids (averaged traces in Fig. 3j; ****P<0.0001, two-way repeatedmeasures ANOVA, n = 5 mice). Panels **h**-**j** show that SFO^{Nos1} neurons do not transmit a teaching signal in a Pavlovian conditioning model. h, Schematic of Pavlovian conditioning model. i, SFO^{Nos1} neurons were not inhibited by cue presentation after 1 week of Pavlovian conditioning (n = 3 mice). **j**, PSTH of SFO^{Nos1} neuron activity and lick rate around the first lick in the first drinking bout either before or after Pavlovian conditioning (n = 3 mice). Panels **k**-**n** demonstrate that SFO^{Nos1} neurons are modulated by rapid anticipatory signals during drinking in the absence of homeostatic need. k, The activity of SFO^{Nos1} neurons was recorded while fully hydrated mice were given ad libitum access to sucrose. l, Representative recording showing modulation of SFO^{Nos1} neurons during sucrose drinking (1 out of 4 mice; red lines, licks). m, PSTH of SFO^{Nos1} neuron activity and lick rate around the first lick in all sucrose drinking bouts (n = 4 mice). n, PSTH of SFO^{Nos1} neuron activity and lick rate around the last lick in all sucrose drinking bouts (n = 4 mice). Values are mean \pm s.e.m. (error bars or shaded area).



Extended Data Figure 6 | **Activation of SFO**^{Nos1} **neurons during eating does not require angiotensin signalling. a**, Experimental design to test whether angiotensin signalling is necessary for prandial thirst. **b**, Angiotensin blockers (INH) did not inhibit eating-induced activation of SFO^{Nos1} neurons or prandial drinking (n = 3 mice). **c**, Angiotensin blockers did not affect food consumption (n.s., two-tailed Student's *t*-test, n = 3 mice). **d**, **e**, ARC^{AgRP} neurons that control hunger are not reciprocally

modulated by eating and drinking. **d**, Schematic of fibre photometry set up for recording the activity of ARC^{AgRP} neurons. Scale bar, 100 μ m. **e**, ARC^{AgRP} neurons were rapidly inhibited when fasted mice were presented with chow, as previously reported²⁷, but were unaffected when dehydrated mice were presented with water (n = 5 mice). Values are mean \pm s.e.m. (error bars or shaded area).



Extended Data Figure 7 | **Silencing of SFO**^{Nos1} **neurons disinhibits feeding. a**, Experimental design to test whether prandial thirst inhibits food intake. **b**, Mice provided simultaneous access to water consumed more food after overnight fasting than mice without simultaneous access to water (**P < 0.01, ****P < 0.0001, two-way ANOVA, n = 10 mice per group), consistent with previous reports that thirst can inhibit hunger in



rats³³. **c**, Experimental design to test whether SFO^{Nos1} neurons mediate inhibition of food intake by prandial thirst. **d**, Silencing of SFO^{Nos1} neurons increased food intake when mice were provided with access to chow without simultaneous access to water after overnight fasting (*****P* < 0.0001, two-way repeated-measures ANOVA, *n* = 5 mice). Values are mean \pm s.e.m. (error bars or shaded area).



Extended Data Figure 8 | **Projection mapping and retrograde tracing from SFO neurons. a**, Schematic of viral strategy for identifying projections from SFO^{Nos1} neurons using a fluorescent synaptophysin fusion protein. **b**, Representative images showing SFO^{Nos1} neuron somas in the SFO and axon terminals in the OVLT, MnPO, PVH and supraoptic nucleus (SON) (1 out of 1 mice; green, GFP; blue, DAPI). Scale bars, Retrograde-labeled neurons

100 µm. c, Schematic of strategy for retrograde tracing from SFO neurons using retrobeads. d, Representative images showing retrobeads injection site in the SFO and retrograde-labelled neurons in the medial septum (MS), OVLT, MnPO, ARC, MnR, dorsal raphe (DR), and locus coeruleus (LC) (1 out of 2 mice; red, rhodamine; blue, DAPI). Scale bars, 100 µm.

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Extended Data Figure 9 | Schematic for convergence of anticipatory and homeostatic signals at SFO^{Nos1} thirst neurons. SFO^{Nos1} neurons monitor the composition of the blood by sensing plasma osmolarity and, via angiotensin, plasma volume and pressure. SFO^{Nos1} neurons predict the future state of the blood by integrating temperature-dependent inputs from the mouth and osmolarity-dependent inputs from the gut during drinking, and angiotensin- and osmolarity-independent inputs from the mouth/gut during eating. Illustration from National Institute of Diabetes and Digestive and Kidney Diseases, US National Institutes of Health.

Extended Data Table 1 | Summary of statistical analyses Figure Sample size (n) Statistical test

Figure	Sample size (n)	Statistical test	Values
1c	5 mice	two-way repeated-measures ANOVA factor one: time-point (baseline, 45 min) factor two: injection (vehicle, 3M NaCl, 3M NaCl + INH) post-hoc correction for multiple comparisons (Holm-Šídák)	time-point: $F(1,4) = 134.96$, $P = 0.0003$ injection: $F(2,8) = 37.79$, $P < 0.0001$ interaction: $F(2,8) = 30.41$, $P = 0.0002$ multiple comparisons: **** $P < 0.0001$
1d	5 mice	two-way repeated-measures ANOVA factor one: time-point (baseline, 45 min) factor two: injection (vehicle, 1M NaCl, 2M mannitol) post-hoc correction for multiple comparisons (Holm-Šídák)	time-point: $F(1,4) = 21.29$, $P = 0.0099$ injection: $F(2,8) = 4.74$, $P = 0.0438$ interaction: $F(2,8) = 13.21$, $P = 0.0029$ multiple comparisons: *** $P < 0.001$
1f	5 mice	two-way repeated-measures ANOVA factor one: time-point (baseline, 5 min) factor two: injection (vehicle, PEG, PEG + INH) post-hoc correction for multiple comparisons (Holm-Šídák)	time-point: $F(1,4) = 55.81$, $P = 0.0017$ injection: $F(2,8) = 26.43$, $P = 0.0003$ interaction: $F(2,8) = 99.26$, $P < 0.0001$ multiple comparisons: **** $P < 0.0001$
1h	5 mice	two-way repeated-measures ANOVA factor one: time-point (baseline, 5 min) factor two: injection (vehicle, isoproterenol, isoproterenol + INH) post-hoc correction for multiple comparisons (Holm-Šídák)	time-point: <i>F</i> (1,4) = 18.59, <i>P</i> = 0.0125 injection: <i>F</i> (2,8) = 36.73, <i>P</i> < 0.0001 interaction: <i>F</i> (2,8) = 18.86, <i>P</i> = 0.0009 multiple comparisons: *** <i>P</i> < 0.001
2a	6 mice	one-way repeated-measures ANOVA factor one: time-point (baseline, water restriction, water re-access) post-hoc correction for multiple comparisons (Holm-Šídák)	time-point: <i>F</i> (2,10) = 77.85, <i>P</i> < 0.0001 multiple comparisons: **** <i>P</i> < 0.0001
2d	5 mice	one-way repeated-measures ANOVA factor one: time-point (baseline, 5 min, 45 min) post-hoc correction for multiple comparisons (Holm-Šídák)	time-point: <i>F</i> (2,8) = 131.40, <i>P</i> < 0.0001 multiple comparisons: **** <i>P</i> < 0.0001
2e	ad lib.: 9 mice –1 min: 4 mice 5 min: 9 mice 45 min: 8 mice	one-way ANOVA factor one: time-point (ad lib., –1 min, 5 min, 45 min) post-hoc correction for multiple comparisons (Holm-Šídák)	time-point: <i>F</i> (3,26) = 6.20, <i>P</i> = 0.0025 multiple comparisons: * <i>P</i> < 0.05
2f	ad lib.: 9 mice –1 min: 4 mice 5 min: 9 mice 45 min: 8 mice	one-way ANOVA factor one: time-point (ad lib., –1 min, 5 min, 45 min) post-hoc correction for multiple comparisons (Holm-Šídák)	time-point: <i>F</i> (3,26) = 4.93, <i>P</i> = 0.0076 multiple comparisons: * <i>P</i> < 0.05
3c	5 mice	one-way repeated-measures ANOVA factor one: time-point (baseline, 5 min, 45 min, 50 min, 90 min) post-hoc correction for multiple comparisons (Holm-Šídák)	time-point: <i>F</i> (4,16) = 64.35, <i>P</i> < 0.0001 multiple comparisons: **P < 0.01, *** <i>P</i> < 0.001, **** <i>P</i> < 0.0001
3d	10 mice per group	one-way ANOVA factor one: time-point (–1 min, 5 min, 45 min, 50 min, 90 min) post-hoc correction for multiple comparisons (Holm-Šídák)	time-point: <i>F</i> (4,45) = 35.80, <i>P</i> < 0.0001 multiple comparisons: *** <i>P</i> < 0.001, **** <i>P</i> < 0.0001
Зf	5 mice	two-tailed paired Student's t-test	t(4) = 7.08, P = 0.0021
3h	5 mice	two-tailed paired Student's t-test	<i>t</i> (4) = 3.61, <i>P</i> = 0.0225
3k	5 mice	one-way repeated-measures ANOVA factor one: water temp. (12°C, 24°C, 36°C) post-hoc correction for multiple comparisons (Holm-Šídák)	water temp.: <i>F</i> (2,8) = 11.71, <i>P</i> = 0.0042 multiple comparisons: ** <i>P</i> < 0.01
4b	10 mice per group	one-way ANOVA factor one: time-point (fed, –1 min, 5 min, 45 min, 50 min) post-hoc correction for multiple comparisons (Holm-Šídák)	time-point: $F(4,45) = 7.05$, $P = 0.0002$ multiple comparisons: * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$
4e	5 mice	two-way repeated-measures ANOVA factor one: time-bin (0–15 min, 15–30 min) factor two: laser ("–laser" trials, "+laser" trials) post-hoc correction for multiple comparisons (Holm-Šídák)	time-bin: $F(1,4) = 5.21$, $P = 0.0845$ laser: $F(1,4) = 2.67$, $P = 0.1777$ interaction: $F(1,4) = 196.00$, $P = 0.0002$ multiple comparisons: ** $P < 0.01$, *** $P < 0.001$
5c	SFO ^{GFP} : 5 mice SFO ^{ChR2} : 8 mice MnPO ^{GFP} : 4 mice PVH ^{GFP} : 4 mice PVH ^{GFP} : 4 mice	two-way ANOVA factor one: brain region (SFO, MnPO, PVH) factor two: vector (GFP, ChR2) post-hoc correction for multiple comparisons (Holm-Šídák)	brain region: $F(2,28) = 27.02$, $P < 0.0001$ vector: $F(1,28) = 97.24$, $P < 0.0001$ interaction: $F(2,28) = 24.11$, $P < 0.0001$ multiple comparisons: **** $P < 0.0001$