Functional Ontogeny of Hypothalamic Agrp Neurons in Neonatal Mouse Behaviors

Graphical Abstract

Highlights

- Isolation from the nest activates Agrp neurons in neonatal mice
- Care and warmth, but not milk, blunts activation of Agrp neurons
- Neonatal Agrp neurons modulate isolation-induced ultrasonic vocalizations
- Agrp neurons increase milk ingestion in 15- but not in 10-day-old mice

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In Brief
Hypothalamic Agrp neurons play a role in offspring-to-caregiver bonding independent of their role in food ingestion.
Functional Ontogeny of Hypothalamic Agrp Neurons in Neonatal Mouse Behaviors

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SUMMARY

Hypothalamic Agrp neurons regulate food ingestion in adult mice. Whether these neurons are functional before animals start to ingest food is unknown. Here, we studied the functional ontogeny of Agrp neurons during breastfeeding using postnatal day 10 mice. In contrast to adult mice, we show that isolation from the nursing nest, not milk deprivation or ingestion, activated Agrp neurons. Non-nutritive suckling and warm temperatures blunted this effect. Using in vivo fiber photometry, neonatal Agrp neurons showed a rapid increase in activity upon isolation from the nest, an effect rapidly diminished following reunion with littermates. Neonates unable to release GABA from Agrp neurons expressed blunted emission of isolation-induced ultrasonic vocalizations. Chemogenetic overactivation of these neurons further increased emission of these ultrasonic vocalizations, but not milk ingestion. We uncovered important functional properties of hypothalamic Agrp neurons during mouse development, suggesting these neurons facilitate offspring-to-caregiver bonding.

INTRODUCTION

Agouti-related peptide (Agrp) neurons in the arcuate nucleus of the hypothalamus serve as a central coordinator to regulate food intake. Ablation of these neurons leads to aphagia in adult mice (Gropp et al., 2005; Luquet et al., 2005), but not in neonates, suggesting that Agrp neurons do not contribute to ingestive behaviors early in development. In support of this view, Agrp neurons show delayed development in rodents (Nilsson et al., 2005; Padilla et al., 2010), when the final maturation of Agrp neuronal circuitry coincides with weaning (Grove and Smith, 2003). However, impairing development of Agrp neurons during the first postnatal week in mice has persistent consequences to metabolism and behavior (Dietrich et al., 2012; Joly-Amado et al., 2012), suggesting an unidentified function for Agrp neurons during early development.

Here, we assessed the functional ontogeny of Agrp neurons in early postnatal development of mice at postnatal day 10 (P10) and during the weaning period during postnatal days 15–21 (P15–P21).

RESULTS

Isolation from the Nursing Nest, Not Nutrient Intake, Activates Agrp Neurons in Neonates

In adult mice, nutrient deprivation activates Agrp neurons (Hahn et al., 1998; Takahashi and Cone, 2005). So, we investigated the extent to which Agrp neurons respond to a lack of nutrients in neonatal mice. To test this, we isolated P10 mice from the nest for 90 min or 8 h to prevent nutrient intake via milk ingestion (Figure 1A). A 90-min time from onset of separation maximizes Fos expression (Barros et al., 2015). Both periods of isolation increased Agrp neuronal activity, as indicated by the increased number of Fos positive Agrp neurons upon isolation (nest: 3.14 ± 0.96%, n = 9; isolation/90 min: 31.19 ± 1.89%, n = 10; isolation/8 h: 46.19 ± 2.04%, n = 3; F2, 19 = 128.6, p < 10−11, one-way ANOVA; Figures 1B–1D).

Because 90 min significantly increased Fos positive Agrp neurons, we tested whether this period of isolation also stimulated milk intake. We measured milk intake by calculating the change in body weight in animals that remained in the nest for 90 min compared to animals isolated for 90 min followed by re-introduction to the nest for an additional 90-min period (Figure 1E). In this protocol, we did not observe significant differences in the body-weight changes between the two experimental conditions (nest: 62.1 ± 15.4 mg, n = 12; reunion: 76.2 ± 14.4 mg, n = 12; t22 = 0.65, p = 0.51, unpaired t test; Figure 1F). Thus, activation of Agrp neurons after 90 min of isolation from the nest in P10 mice is not significant to increase milk intake. Moreover, the increased activation of Agrp neurons does not seem to arise from a generalized stress response, as corticosterone levels did not increase after isolation (nest: 9.96 ± 1.85 ng/mL, n = 7; isolation: 12.40 ± 1.85 ng/mL, n = 11; t16 = 0.88, p = 0.39, unpaired t test; Figure 1G), and testing pups in the presence of a predator odor for 90 min did not alter Fos labeling in Agrp neurons (nest: 1.19 ± 0.92%, n = 2; isolation: 23.82 ± 4.78%, 178–179). This finding suggests that activation of these neurons is not related to a generalized stress response.
Figure 1. Fos Labels Agrp Neuron Activation in P10 Mice upon Social Isolation

(A) P10 mice were socially isolated for 90 min or 8 h.

(B) Agrp neurons positive for Fos immunoreactivity (nest, n = 9; isolation/90 min, n = 10; isolation/8 h, n = 3, one-way ANOVA, p < 10^-11).

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To evaluate whether the activation of Agrp neurons upon isolation from nest is due to caloric deprivation, we isolated P10 mice while orally infusing bovine or mouse derived milk (Figure 1K). We confirmed milk ingestion by adding a colored dye to the milk to verify its presence in the stomach (Figure 1L). Surprisingly, our results showed no significant difference between the activation of Agrp neurons between the groups that received an oral infusion of milk and the control group that received a passive oral probe with no milk infusion (nest: 1.51 ± 0.65%, n = 5; isolation and no infusion: 27.88 ± 2.39%, n = 12; isolation and mouse milk: 30.73 ± 1.93%, n = 8; isolation and bovine milk: 24.24 ± 3.14%, n = 8; F3, 29 = 19.38, p < 10^-6, one-way ANOVA; Figures 1M–1O).

Next, we tested the hypothesis that Agrp neuron activity in P10 mice increases upon separation in anticipation of future nutrient deprivation. To this end, we investigated the extent to which isolation from the nest activates Agrp neurons when dissociating milk intake and the nest. P7 mice were housed with a foster non-lactating dam and were manually fed milk by an investigator (Figure 1P). All neonates quickly developed locomotor activity toward the investigator at feeding onset, which suggests a learned association that the milk source was outside the home nest. We then assessed activation of Agrp neurons at P10 at either isolation or return to the home cage with the foster dam for a period of 90 min after all pups were previously fed with equal volumes of milk. Similar to our previous experiments, isolation from the home nest strongly increased the number of Fos positive Agrp neurons compared to mice that returned to the home cage (foster dam: 15.59 ± 3.26%, n = 6; isolation: 64.39 ± 3.94%, n = 8; t12 = 8.06, p < 10^-5, unpaired t test; Figures 1Q and 1R). Taken together, these results suggest that activation of Agrp neurons in P10 mice following isolation from the nest does not require milk deprivation or anticipation of milk deprivation, which stands in contrast to adult mice.

Non-nutritive Suckling and Thermal Support Blunt Agrp Neuron Activation in Neonates

Our next goal was to evaluate the relative importance of different components of the nest environment and mother-infant interaction that contribute to Agrp neuron activation in P10 mice after isolation from the nest. In the nursing nest, pups receive care from the dam. An important feature of maternal care is neonatal attachment to the mother’s nipple and suckling. So, we investigated the extent to which suckling alters the activation of Agrp neurons. We fostered P10 mice with non-lactating dams, non-lactating dams with protruded nipples, and lactating dams (Figure 2A). In all cases, foster dams promptly retrieved the pups and placed them in the nest. All foster dams displayed maternal behaviors, such as grooming and licking and arched-back “nursing” of pups, as expected. Interestingly, all foster dams blunted the activation of Agrp neurons in P10 mice compared to isolated pups (nest: 2.10 ± 0.21%, n = 6; isolation: 26.47 ± 1.57%, n = 6; non-lactating foster dam: 16.59 ± 1.64%, n = 7; non-lactating foster dam with protruded nipples: 8.66 ± 1.26%, n = 6; lactating foster dam: 10.29 ± 0.65%, n = 4; F4, 24 = 50.29, p < 10^-10, one-way ANOVA; Figures 2B and 2C). Attachment of pups to the foster dam’s nipples further decreased the number of Fos-labeled Agrp neurons compared to pups placed with a foster dam with non-protruded nipples to prevent nipple attachment (Figures 2B and 2C). The effect of nipple attachment was irrespective of milk availability, as the expression of Fos in pups showed a similar magnitude when placed with lactating and non-lactating foster dams with protruded nipples (Figures 2B and 2C). Overall, activation of Agrp neurons in P10 mice is blunted by non-nutritive suckling, an important component of maternal care, and is not further reduced by availability of milk in the dam’s nipples.
Figure 2. Warm Temperatures Blunt Activation of Agrp Neurons in P10 Mice

(A) Study design: P10 mice were either fostered with non-lactating dams, with non-lactating dams with protruded nipples, or with lactating dams for 90 min. A control group was not manipulated (nest) and a second group was isolated (isolation).

(B) Quantification of Agrp neurons positive for Fos (nest, n = 6; isolation, n = 6; non-lactating foster dam, n = 7; non-lactating foster dam with protruded nipples, n = 6; lactating foster dam, n = 4, one-way ANOVA, p < 10^{-10}).

(C) Tukey-Kramer’s multiple comparisons test of the difference between means (95% confidence intervals from B), representing effect sizes.

(D) Thermo-photography of the nursing nest. Lactating dam is on the top of P10 offspring. Nest temperature is approximately 34–36°C.

(E) Offspring raised at room temperature (RT) were isolated either at room temperature, at thermoneutrality (TN, in a climate chamber set to 35°C) or at room temperature with a thermal support (TS) irradiating heat from underneath the cage (≈35°C).

(F) Quantification of Agrp neurons positive for Fos (nest and room temperature, n = 5; isolation and room temperature, n = 7; isolation and thermoneutrality, n = 10; isolation and thermal support, n = 8, one-way ANOVA, p = 10^{-10}).

(G) Tukey-Kramer’s multiple comparisons test of the difference between means (95% confidence intervals from F) representing effect sizes.

(H) Offspring raised at thermoneutrality (climate chamber at 35°C) were isolated for 90 min at room temperature or thermoneutrality.

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The nursing nest provides critical thermal insulation, which reduces heat loss from neonates that have not fully developed homeostatic mechanisms for thermoregulation. Dams contribute to thermal insulation by building a nest and skin-to-skin contact with pups (Figure 2D). To test the effects of thermal insulation on the activation of Agrp neurons, we isolated P10 mice at room temperature or at a thermoneutral temperature (= 35°C; Figure 2E). Because temperature exchanges in the nest occur by skin-to-skin contact, we included an additional control group, in which we provided thermal support by irradiating heat (=35°C) from underneath (Figure 2E). 90 min of isolation at thermoneutrality or with thermal support strongly suppressed activation of Agrp neurons in P10 mice as assayed by Fos labeling compared with pups isolated at room temperature (nest: 0.77 ± 0.39%, n = 5; isolation and room temperature: 22.19 ± 2.53%, n = 7; isolation and thermoneutrality: 5.90 ± 0.86%, n = 10; isolation and thermal support: 2.75 ± 0.51%, n = 8; F2, 26 = 47.31, p = 10^-10, one-way ANOVA; Figures 2F and 2G). Thus, Agrp neurons in P10 mice respond to the withdrawal of thermal insulation when isolated from the nursing nest. This factor holds primary importance for the responses of these neurons to isolation.

Next, we tested the extent to which the response of Agrp neurons in P10 mice to the withdrawal of thermal insulation was dependent on prior experience with drops in ambient temperature. To prevent mice from experiencing ambient temperatures lower than nest temperatures, we repeated the experiments in animals born and raised in a thermoneutral environment (Figure 2H). P10 mice raised at thermoneutrality showed increased Fos-labeled Agrp neurons when isolated at room temperature but not at thermoneutrality (nest and thermoneutrality: 0.62 ± 0.33%, n = 2; isolation and room temperature: 27.94 ± 3.13%, n = 3; isolation and thermoneutrality: 6.16 ± 1.54%, n = 3; F2, 5 = 37.91, p = 0.001, one-way ANOVA; Figures 2I and 2J). Thus, the response of Agrp neurons to withdrawal of thermal insulation in P10 mice does not require previous experiences with drops in ambient temperature.

**Neonatal Agrp Neurons Undergo Rapid Activity Changes**

In the previous experiments, we could not elucidate the temporal dynamics of physiological activation of Agrp neurons. For example, Agrp neurons after isolation (Figure 3A) could slowly increase their activity similar to a homeostat (Figure 3B). Alternatively, these neurons could rapidly respond to isolation (Figure 3C) similar to an alarm and reflexive system. A third alternative suggests that Agrp neurons could show delayed activation (Figure 3D), suggesting a thresholding mechanism triggers these neurons in neonates.

To better understand the natural activity dynamics of Agrp neurons early in postnatal development, we injected an adenovirus encoding [GCaMP7s in a Cre-dependent manner in newborn AgrpCre/Cre mice (Figures 3E–G) (Dana et al., 2018). We then used fiber photometry to measure calcium transients originating from Agrp neurons upon isolation-reunion in P13–14 pups (Figure 3A). We found that pup isolation from the nest increased activity of Agrp neurons that occurred within seconds (Figures 3H and 3I) and persisted throughout the isolation period (10 min). After this initial separation, reunion of the isolated animal with the litter immediately decreased the activity of Agrp neurons (Figures 3H and 3I). The suppression of Agrp neuronal activity was robust and rapid, normalizing the detected signal to pre-isolation levels in less than 30 s (Figures 3H and 3I).

All animals tested showed this response to isolation-reunion, suggesting a general model in which Agrp neurons in neonates rapidly respond to disruptions in the nest conditions (Figure 3C).

**Neonatal Agrp Neurons Modulate the Emission of Ultrasonic Vocalizations**

In most neonatal mammals, including mice, disruptions in the nest condition lead to infant vocalization (Hofer, 1994). In mice and rats, neonates emit ultrasonic vocalizations (USVs) when separated from the dam (Noirot, 1966, 1968; Zippeius and Schleidt, 1956) (Figures 4A and 4B). We investigated whether activation of Agrp neurons in neonates upon isolation from the nest could modulate emission of USVs.

First, we confirmed that isolation from the nest induced vocal behavior in P10 mice (Figure 4A). We then investigated whether isolation at thermoneutrality would influence USVs, since these conditions blunt activation of Agrp neurons upon isolation (Figure 2). Analysis of vocal behavior showed a rapid increase in USV emission upon isolation from the nest (Figures 4B–4E), an effect that blunted at thermoneutrality (nest, n = 3; isolated and room temperature, n = 16; isolated and thermoneutrality, n = 4; F2, 20 = 18.08, p < 10^-4, one-way ANOVA; Figures 4C–4E). Thus, vocal behavior dynamics in neonatal mice follow the dynamics of Agrp neuron activation upon isolation in these experimental conditions.

To test the extent to which Agrp neurons contribute to vocal behavior of P10 mice upon isolation, we tested animals lacking the transmitters released by Agrp neurons (NPY and GABA) (Hahn et al., 1998; Horvath et al., 1997). We recorded the emission of USVs in NpyKO and AgrpVgat-KO mice and their littermate controls following 10 minutes isolation in P10 mice. Animals lacking NPY exhibited a similar number of USVs after isolation compared to controls (control: 343.4 ± 56.8 USVs, n = 12; NpyKO: 384.5 ± 69.3 USVs, n = 17; NpyKO/+/: 348.0 ± 91.7 USVs, n = 7; p = 0.74, Kruskal-Wallis [KW] test; Figure 4F). In contrast, AgrpVgat-KO mice had a significant decrease in emission of USVs upon isolation (control: 321.0 ± 50.4 USVs, n = 10; AgrpVgat-KO: 57.7 ± 14.6 USVs, n = 14; U = 2, p < 10^-5, Mann-Whitney test; Figures 4G and 4H).

We further analyzed the spectro-temporal characteristics of 3,427 USVs from control mice and 786 USVs from AgrpVgat-KO mice. We characterized individual USVs by changes in the transmitters released by Agrp neurons (NPY and GABA) and the transmitters released by Agrp neurons (NPY and GABA) and the transmitters released by Agrp neurons (NPY and GABA). We found that control mice showed a significant decrease in emission of USVs upon isolation (control: 321.0 ± 50.4 USVs, n = 10; AgrpVgat-KO: 57.7 ± 14.6 USVs, n = 14; U = 2, p < 10^-5, Mann-Whitney test; Figures 4G and 4H).
spectro-temporal characteristics, such as duration, frequency, and bandwidth. Compared to control mice, USVs from Agrp$^{Vgat-KO}$ mice decreased in duration of 9.5 ms (control: 37.84 ± 0.45 ms; Agrp$^{Vgat-KO}$: 28.31 ± 0.84 ms; $D = 0.176, p = 10^{-10}$, Kolmogorov-Smirnov [KS] test; Figure 4I), in mean frequency of 1.9 kHz (control: 82.20 ± 0.28 kHz; Agrp$^{Vgat-KO}$: 80.24 ± 0.55 kHz; $D = 0.10, p = 10^{-15}$, KS test; Figure 4J), and in bandwidth of 5.6 kHz (control: 22.86 ± 0.35 kHz; Agrp$^{Vgat-KO}$: 17.20 ± 0.67 kHz; $D = 0.17, p = 10^{-15}$, KS test; Figure 4K). We also found an overall decrease in the number of vocalizations across most USV categories (Figures 4L–4X) (Grimsley et al., 2011). However, we observed an increase in the incidence of “short” vocalizations, when analyzing the relative frequency of USV categories (Figure 4Y). This syllable represents the simplest form of vocalization by neonatal mice based on spectro-temporal characteristics (Figures 4L–4V). Thus, lacking GABA release from Agrp neurons, P10 Agrp$^{Vgat-KO}$ mice led to fewer and simpler USVs compared to control animals. Taken together, these findings indicate Agrp neurons are critically positioned to modulate the emission of USVs in neonatal mice.

**Chemogenetic Activation of Agrp Neurons Increases USV Emission**

We further tested whether chemogenetic activation of Agrp neurons could modulate emission of USV in isolated P10 pups using Agrp$^{Trpv1}$ mice (Dietrich et al., 2015; Ruan et al., 2014; Arenkiel et al., 2008; Güler et al., 2012) (Figure 5A; Figure S1). Subcutaneous injection of capsaicin (10 mg/kg) in Agrp$^{Trpv1}$ mice robustly activated Agrp neurons in young pups (Figure 5B; n = 5 mice per group; $U = 0, p = 0.004$, Mann-Whitney test). Chemogenetic activation of Agrp neurons using the Agrp$^{Trpv1}$ animal model induced a 61% increase in USV emission in P10 mice (control: 686.7 ± 50.22 USVs, n = 32; Agrp$^{Trpv1}$: 1040.0 ± 66.56 USVs, n = 24; $t_{51} = 4.318, p < 0.0001; 2$-tailed unpaired $t$ test; Figures 5C–5E).

Figure 3. Rapid Dynamics of Agrp Neuronal Activity in Mice during Early Development

(A) Experimental model of isolation and reunion in neonates, which activates Agrp neurons after 90 min. Three theoretical models of activity changes of these neurons are illustrated in (B)–(D).

(B) Model A: activity of Agrp neurons gradually increases during the 90-min isolation.

(C) Model B: activity of Agrp neurons rapidly increases upon isolation.

(D) Model C: activity of Agrp neurons increases in isolation after a delay.

(E) In newborn Agrp$^{Cre/Cre}$ mice, an adeno-associated virus was injected in the arcuate nucleus of the hypothalamus to express jGCaMP7s in Agrp neurons (AAV-CAG-Flex-jGCaMP7s).

(F) Preweaning mouse connected to an optic fiber and its dam.

(G) Expression of jGCaMP7s in Agrp neurons of a P14 mouse.

(H) Z score of Agrp neuronal activity in P13–14 mice. Baseline was recorded for 1 min and then pups were isolated for 10 min. Subsequently, pups were reunited with the litter. Plot represents mean ± SEM (n = 7 animals).

(I) Heat plot representing individual responses to isolation and reunion.
Figure 4. Agrp Neurons in P10 Mice Modulate Emission of Ultrasonic Vocalizations via GABA Release
(A) Experimental model of isolation in P10 mice at room temperature (RT) or at thermoneutral conditions (thermoneutrality; climate chamber set at 35°C).
(B) Representative spectrogram of ultrasonic vocalizations (USVs) in P10 mice recorded in isolation.
(C) Number of USVs in five-minute bins during isolation at room temperature or thermoneutrality and control group recorded in the nest.
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We also analyzed the spectro-temporal characteristics of a total of 21,019 USVs from control mice and 24,976 vocalizations from Agrp<sup>Trpv1</sup> mice. Upon activation of Agrp neurons, the USV duration decreased by 4 ms (control: 51.42 ± 0.23 ms; Agrp<sup>Trpv1</sup>: 47.09 ± 0.19 ms; *D* = 0.077, *p* = 10<sup>-15</sup>, KS test; Figure 5F), the mean frequency increased by 3 kHz (control: 77.50 ± 0.08 kHz; Agrp<sup>Trpv1</sup>: 80.96 ± 0.07 kHz; *D* = 0.12, *p* = 10<sup>-15</sup>, KS test; Figure 5G), and the bandwidth increased by 2 kHz (control: 27.31 ± 0.13 kHz; Agrp<sup>Trpv1</sup>: 29.84 ± 0.12 kHz; *D* = 0.07, *p* = 10<sup>-15</sup>, KS test; Figure 5H). The selected categories of USVs induced by activation of Agrp neurons (Figure 5I) showed a higher complexity compared to those suppressed in Agrp<sup>Vgat-KO</sup> mice (Figure 4).

Interestingly, when we analyzed the relative frequency of USV categories, we found a selective increase in the frequency of “chevrons” upon chemogenetic activation of Agrp neurons (Figures 5J and 4N). To further corroborate these findings, we used a second software tool MUPET to classify vocalizations based on their shape (Van Segbroeck et al., 2017). We found six clusters of vocalizations that showed a more than 2.5-fold increase upon activation of Agrp neurons (Figures 5K and 5L). These clusters were very similar to each other, resembling our previous analysis (Figures 4G and 4H). We also found that activation of Agrp neurons suppressed vocalizations in cluster 39 (Figures 5K and 5M). The USVs in cluster 39 were simpler in shape, resembling vocalizations predominantly emitted by Agrp<sup>Vgat-KO</sup> pups (Figure 4). In contrast to mice lacking GABA release by Agrp neurons, chemogenetic activation of these neurons stimulated USV emission at a higher rate and with higher complexity. Together, these results strongly suggest a model in which neonatal Agrp neurons are rapidly activated upon isolation from the nest, which modulates USV emission, presumably to attract the dam.

### Chemogenetic Activation of Neonatal Agrp Neurons Increases Odds for Nipple Attachment

In young pups, contact with the dam is critical for suckling behavior and ingestion of breast milk. Thus, we next devised behavior experiments to test whether activation of Agrp neurons in pups would drive behaviors toward the dam, including exploratory activity, suckling, and milk intake. We eliminated active participation by the dam as the driver of these behaviors by examining the behavior of P10 mice toward anesthetized dams (Figure 6A). In anesthetized dams, milk ejection decreases considerably (Lincoln et al., 1973), so suckling under these conditions is considered non-nutritive. Indeed, we did not observe the stretching reflex in pups that suckled during our experiments, which is a pathognomonic sign of milk ejection and ingestion (Vorherr et al., 1967).

Activation of Agrp neurons in P10 mice increased the total number of pups that attached to the dam’s nipples (control: 9 out of 22; Agrp<sup>Trpv1</sup>: 13 out of 15; *p* = 0.005, chi-square test; Figure 6B) and increased the distance traveled in the testing chamber (control: 0.75 ± 0.14 m, *n* = 22; Agrp<sup>Trpv1</sup>: 1.20 ± 0.21 m, *n* = 15; *U* = 94, *p* = 0.027, Mann-Whitney test; Figures 6C and 6D). We then compared nipple attachment behavior of P10 mice, excluding animals that did not attach to the dam’s nipples from post hoc analysis. Chemogenetic activation of Agrp neurons did not change the frequency (control: 3.11 ± 1.23; Agrp<sup>Trpv1</sup>: 3.61 ± 0.83, *U* = 51, *p* = 0.63, Mann-Whitney test; Figure 6E), latency (control: 279.9 ± 57.4 s; Agrp<sup>Trpv1</sup>: 485.4 ± 86.2 s; *U* = 35, *p* = 0.12, Mann-Whitney test; Figure 6F), or the duration of nipple attachment (control: 719.4 ± 128.0 s; Agrp<sup>Trpv1</sup>: 529.8 ± 99.86 s; *U* = 39, *p* = 0.20, Mann-Whitney test; Figure 6G). Thus, while chemogenetic activation of Agrp neurons in P10 mice increased the probability of attaching to the dam’s nipples, it did not change the observable microstructure of nipple attachment behavior.
Figure 5. Activation of Agrp Neurons in P10 Mice Increases USV Emission and Alters the Dam’s Behavior

(A) Generation of Agrp<sup>Trpv1<sup>−/−</sup> mice.

(B) Fos in the arcuate nucleus of P15 Agrp<sup>Trpv1<sup>−/−</sup> mice upon injection of capsaicin (10 mg/kg, s.c.; n = 5 mice per group). Scale bar corresponds to 50 μm. Bars and symbols represent mean ± SEM.

(C) Raster plots show USV in isolated P10 controls (n = 32) and Agrp<sup>Trpv1<sup>−/−</sup> mice (n = 24). A tick represents a USV.

(D) Related to (C), number of USVs (in 5-min bins).

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Next, we investigated the influence of the environmental temperature on nipple attachment behavior (Figure 6H). Thermal support from underneath the testing chamber completely suppressed nipple attachment behavior in P10 control mice (0 out of 11 pups tested attached), while 4 out of 10 pups with activated Agrp neurons attached to the dam’s nipples (p = 0.019, chi-square test; Figure 6I). Interestingly, the arousal response of P10 mice upon activation of Agrp neurons was intact, as measured by the distance traveled during the test (control: 0.30 ± 0.07 m, n = 11; AgrpTrpv1: 1.03 ± 0.19 m, n = 10; U = 13, p = 0.002, Mann-Whitney test; Figures 6J and 6K). When we analyzed the different components of nipple attachment behavior of the four P10 mice that successfully attached, we found the frequency of attachments was largely suppressed with mice only attaching once during the test (Figure 6L). The latency to attach (386.5 ± 49.62 s; Figure 6M) was similar to the other experimental conditions (Figures 6F and 6S). The duration animals remained attached to the nipples (787.9 ± 48.20 s; Figure 6N) was within the range of our other experiments (Figures 6G and 6T). We conclude providing warmth did not blunt the arousal response after activating Agrp neurons but did suppress nipple attachment behavior of P10 mice. Taken together, our behavioral experiments further support the importance of a thermal stimulus in neonates to modulate the functional properties of Agrp neurons.

We then assessed whether a lactating anesthetized dam would alter attachment by P10 mice. Oxytocin triggers the milk ejection reflex following nipple stimulation by the pups during suckling (Lincoln and Paisley, 1982). To facilitate milk ejection, we injected a group of anesthetized dams with oxytocin immediately before testing each pup (Figure 6O) (Singh and Hofer, 1978; Vorherr et al., 1967). Similar to non-lactating dams, activation of Agrp neurons in P10 mice increased the number of mice that attached to the dam’s nipples (p = 0.019, chi-square test; Figure 6I). Interestingly, upon activation of Agrp neurons, P10 mice ingested a lower amount of milk than controls during the test (control: 91.2 ± 19.9 mg; AgrpTrpv1: 43.7 ± 10.0 mg; pups that did not attach: −10.0 ± 2.9 mg, n = 12; F2,30 = 19.35, p < 10−5, one-way ANOVA; Figures 6U and 6V). These results suggest that activated Agrp neurons increase dam-seeking behavior but not necessarily increase ingestion of milk.

Chemogenetic Activation of Agrp Neurons Increases Ingestive Behaviors in P15 Mice

Mice rapidly transition from breastfeeding to independent feeding during weaning period. At approximately P15, mice begin experimenting with food sources but still rely on breastfeeding for nutrition (Hammond et al., 1996). We next examined the behavior of P15 mice toward the dam to investigate the ontogeny of Agrp neuron function. All tested P15 mice attached to the nipples of the anesthetized dam regardless of chemogenetic activation of Agrp neurons (n = 19 mice per group; Figure 7A). Activating Agrp neurons did not significantly change the distance traveled in the testing chamber (control: 2.03 ± 0.34 m; AgrpTrpv1: 2.20 ± 0.22 m; U = 145, p = 0.31, Mann-Whitney test; Figure 7B). In contrast to P10 mice, chemogenetic activation of Agrp neurons in P15 mice showed a striking increase in the number of attachments to the dam’s nipples compared to control (3.68 ± 1.02; AgrpTrpv1: 11.74 ± 1.80; U = 72.5, p = 0.001; Mann-Whitney test; Figures 7C and 7D). We observed no statistical difference in the latency of the first attachment (control: 226.6 ± 50.8 s; AgrpTrpv1: 123.0 ± 16.92 s; U = 116, p = 0.06; Mann-Whitney test; Figure 7E) or the total duration of nipple attachment (control: 711.6 ± 85.2 s; AgrpTrpv1: 798.9 ± 79.02 s; U = 143, p = 0.28; Mann-Whitney test; Figure 7F). Since P15 mice displayed numerous nipple attachments (Figure 7D), we could track and quantify the number of nipples

(E) Related to (C) and (D), total number of USVs in the 20 min after activating Agrp neurons in isolated pups (control, n = 32; AgrpTrpv1, n = 24; p value calculated using a 2-tailed unpaired t test).
(F–H) Violin plots representing the distribution of USV characteristics in control (n = 22,168 USVs) and AgrpTrpv1 mice (n = 24,971 USVs) in (F) duration, (G) mean frequency of the main component of the USV, and (H) the bandwidth; p values calculated using the Kolmogorov-Smirnov test.
(I) Distribution of absolute counts of each vocal call type.
(J–L) Similar to (I), but vocal types normalized to total counts. In (J) and (L), multiple t tests with p values corrected for multiple comparisons using the Holm-Sidak method.
(K) Distribution of the fold change (number of USVs from AgrpTrpv1 mice related to number of USVs from control mice) in each of the 60 output clusters analyzed by MUPET (see STAR Methods). Shown in blue are 6 clusters highly enriched in USVs from AgrpTrpv1 mice. Shown in green is 1 cluster enriched in USVs from controls.
(M) Related to (K), image represents the 6 clusters enriched in USVs from AgrpTrpv1 mice.
(N) Maternal preference test (MPT).
(O) Representative tracking (in blue) of a dam in the preference stage.
(P) Preference index (in seconds) during MPT (n = 25 pairs; p value calculated using a 2-tailed paired t test).
In (E), symbols represent mean ± SEM. In (B), (I), and (J), bars represent mean ± SEM. In (F)–(H), data distribution is plotted. In (E) and (P), symbols represent individual data. In (P), black lines indicate mean ± SEM. See also Figure S1.
Figure 6. Agrp Neuron Activation and Suckling Behavior in P10 Mice

(A) Chamber to assay suckling behavior in mice.

(B–G) Quantification of suckling behavior in P10 mice tested with an anesthetized, non-lactating dam. In (B), proportion of mice displaying nipple attachment. In (C), total distance traveled. In (D), tracking of locomotor activity (bottom: starting point; top: anesthetized dam). In (E)–(G), data only considering mice that attached to nipples. In (E), number of nipple attachments. In (F), latency to the first attachment. In (G), total time attached to nipples.

(H) Infrared (IR) light

(I) IR Camera #2

(J) IR Camera #1

(K) IR Camera #2

(L) IR Camera #1

(legend continued on next page)
explored to measure nipple-shifting behavior (Figure 7G) (Cramer et al., 1980). Chemogenetic activation of Agrp neurons in P15 mice increased the total number of different nipples explored during the test (control: 2.68 ± 0.57 nipples; AgrpTrpv1: 5.21 ± 0.62 nipples; U = 92.5, p = 0.007; Mann-Whitney test; Figure 7H). In these experiments, mice did not show a nipple preference (n = 38 mice; p = 0.42, one-way ANOVA; Figures 7I and 7J), which is a phenomenon observed in other species (Erwin et al., 1975; Hudson et al., 2009; Tomaszynski et al., 1998).

We repeated the above experiments using anesthetized dams injected with oxytocin during a 10-min test. In control mice, 50% of P15 mice attached to the dam’s nipples (4 out of 8 mice), while 91% of AgrpTrpv1 mice displayed the same behavior (11 out of 12 mice; p = 0.035, chi-square test; Figure 7K). Activating Agrp neurons did not significantly change the distance traveled in the testing chamber (control: 1.13 ± 0.26 m; AgrpTrpv1: 1.56 ± 0.21 m; U = 31, p = 0.20, Mann-Whitney test; Figure 7L) but strongly increased the number of nipple attachments (control: 2.00 ± 0.70; AgrpTrpv1: 9.90 ± 1.82; U = 2.5, p = 0.007, Mann-Whitney test; Figure 7M). The latency to the first nipple attachment (control: 162.0 ± 33.1 s; AgrpTrpv1: 119.9 ± 27.1 s; U = 14, p = 0.34, Mann-Whitney test; Figure 7N) and the total duration of nipple attachment (control: 332.2 ± 11.9 s; AgrpTrpv1: 388.0 ± 34.7 s; U = 20, p = 0.85, Mann-Whitney test; Figure 7O) were not changed upon chemogenetic activation of Agrp neurons. In contrast to P10 mice, chemogenetic activation of Agrp neurons in P15 mice did significantly increase milk intake in mice that attached to the nipples of lactating dams (control: 17.2 ± 26.1 mg; AgrpTrpv1: 109.5 ± 38.7 mg; U = 7, p = 0.02, Mann-Whitney test; Figure 7P).

We also tested the extent to which activation of Agrp neurons induces ingestion of solid food during the weaning period. We did not observe changes in food intake in P15 mice, but we observed increased food intake in P18 and P21 mice upon chemogenetic activation of Agrp neurons (Figure 7Q). Similarly, we only found changes in body weight during the feeding test in P21 mice (Figure 7R). Together, this set of behavioral experiments suggest that Agrp neurons in P10 mice are not proximally involved in milk intake or signaling milk ingestion. We propose that these functional properties of Agrp neurons rapidly change (or appear) as mice approach weaning age.

**DISCUSSION**

Overall, our results reveal functional properties of Agrp neurons in neonatal mice. These insights demonstrate developmental differences that emerge during ontogeny, so studying any complex system may remain incomplete without assessing its developmental properties (Tinbergen, 1963).

Our experiments unexpectedly revealed that Agrp neurons are functional during the first 2 postnatal weeks in mice despite their immature characteristics (Nilsson et al., 2005; Padilla et al., 2010). We showed that Agrp neurons in P10 mice did not respond to milk intake and their activation did not directly increase milk intake. Conversely, non-nutritive suckling and thermal insulation were key factors modulating the activity of Agrp neurons. These results are compatible with the physiology of breastfeeding in neonatal mammals. During breastfeeding, mice, like most other mammals, do not receive continuous milk ejection. Milk ejection remains under control of a neuroendocrine reflex (Cross and Harris, 1952) and occurs at random intervals, as demonstrated in rats (Lincoln et al., 1973). In spite of milk ejection patterns, neonatal rats stay attached to the dam’s nipple for at least 12 h a day in the first days of life (Lincoln et al., 1973). In fact, homeostatic sensing of milk deprivation to modulate nutritive sucking behavior only develops later as shown in laboratory rodents (Ellis et al., 1984; Hall and Rosenblatt, 1978; Kenny et al., 1979). Thus, these studies strongly imply that nipple attachment serves as a stimulus for more than milk ejection. In fact, neonatal rodents develop filial huddling to dams triggered by thermotactile stimulation rather than provision of milk (Alberts, 2007; Alberts and May, 1984). Similar to rodents, neonatal monkeys prefer a cloth mother that provides thermal and tactile stimuli to a wired mother with a nursing bottle (Harlow, 1958), establishing that maternal comfort has a superior importance compared to milk intake in driving neonatal affectional responses. We posit that Agrp neurons of neonates drive this milk-independent encoding of the offspring-to-caregiver bond.

In our studies, thermal insulation was the primary factor modulating the activation of Agrp neurons in neonates following isolation from the nest. Notably, experiencing previous thermal challenges was not significant to activate neonatal Agrp neurons following isolation from the nest, which suggests an “innate” property of Agrp neurons. Intriguingly, foster dams also provide thermal insulation for the neonates, but they do not suppress activation of Agrp neurons after isolation from...
Figure 7. Ontogeny of Ingestive Behaviors in Mice

(A–J) Suckling behavior in P15 mice using anesthetized non-lactating dams during a 20-min test. (A) All pups attached to the dam’s nipples during the test. (B) Total distance traveled. (C) Raster plot, ticks indicate nipple attachment. (D) Number of nipple attachments. (E) Latency to the first nipple attachment. (F) Total time attached to nipples. (G) Representative tracking data of nipple attachment. (H) Number of nipples explored. (I) Illustration of the location and number of nipples in the dam and (J) nipple preference as measured by the number of nipple attachments per nipple row in P15 mice (n = 38; both control and AgrpTrpv1 mice were included).

(K–O) Suckling behavior in P15 mice using anesthetized lactating dams during a 10-min test. (K) Proportion of P15 mice that attached to the nipples of lactating dams. In (L), total distance traveled during the test. In (M)–(O), only showing mice that attached to nipples. In (M), the number of nipple attachments is shown. In (N), latency to the first attachment is shown. In (O), total time attached to nipples is shown.

(P) Delta body weight after the suckling assay (control, n = 4; AgrpTrpv1 n = 11).

(Q) Chow intake during 30 min after activation of Agrp neurons in mice at 15, 18, and 21 days of age.

(R) Delta body weight in the same animals as in (Q).

In (A) and (K), statistical analysis performed using the chi-square test (2-tailed). In yellow, proportion of mice that attached to the dam’s nipples. In gray, mice that did not attach. In (B), (D)–(F), (H), and (L)–(P), violin plots represent the distribution of the data. Symbols represent individual values. Statistical analysis performed using Mann-Whitney test. In (Q) and (R), differences tested using unpaired t test with Welch’s correction (unequal SDs) are shown. p values provided in the panels when statistically significant. Bars and symbols (in J, Q, and R) represent mean ± SEM.
the nest to the same degree as thermal support. This observation suggests that neonates integrate several sensory signatures coupled to the nest environment, such as sensory cues from the dam, siblings, home-nest odors, or from all these sources, to create and build expectations of the external world. We unexpectedly propose that activity of Agrp neurons partially encodes this information. Future studies should address the exact nature of this information and the combination of sensory modalities needed to modulate the activity of neonatal Agrp neurons.

Our results provide further evidence to the proposal that Agrp neurons serve as motivational drivers in the mammalian brain. In adult mice, Agrp neurons may encode negative valence (Betley et al., 2015). Under this proposal, these neurons when active would generate an unpleasant state of hunger leading the adult mouse to engage in behaviors to eat and suppress this state (Betley et al., 2015). Our results support this functional property of Agrp neurons in encoding an overall negative state beginning in early development. Clearly, isolation from the nursing nest serves as a negative stimulus for the neonate, which triggers a rapid activation of Agrp neurons and emission of vocalizations. Conversely, reunion then serves as a positive stimulus for the isolated neonate, which immediately suppresses the activity of Agrp neurons. By studying the functional ontogeny of Agrp neurons, our results support a model by which Agrp neurons generate a motivational drive to suppress an overall negative state.

In our studies, we could not find increases in corticosterone levels upon isolation or an enhanced activation of Agrp neurons (as labeled by Fos) in the presence of predator odor. Despite these negative results, we cannot rule out the possibility that Agrp neurons early in life are responsive to general stressors.

Upon parental separation, some infant mammals and birds respond with protest, despair, and emit sounds to attract them (Hofer, 1994). Our results demonstrated that neonatal Agpn neurons are a critical component of the circuit modulating vocal behavioral response in mice. These findings provide further insight into the underlying neural pathways subserving neonatal vocal behavior (Curry et al., 2013; Mosienko et al., 2015; Winslow et al., 2000). Investigating downstream circuits linking Agpn neurons to brain regions involved in the emission of USVs (Arriaga and Jarvis, 2013; Arriaga et al., 2012) will reveal how the neonatal brain encodes sensory information and internal state-dependent variables to generate behavioral responses (Boulanger-Bertolus et al., 2017; Hofer, 1996).

Our results have a broad impact to understand the functional ontogeny of hypothalamic neurons and the importance of the neonatal period in brain and behavior development. Additionally, these studies establish a mechanistic substrate underlying infant-caregiver interaction, which suggests an initial population of neurons underlying the long-sought nature of this social bond in mammals (Harlow, 1958; Lewis et al., 2007).

STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:

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AUTHOR CONTRIBUTIONS

M.R.Z. performed the experiments, designed the studies, analyzed the data, and helped write the manuscript. A.H.O.F. developed a tool for ultrasonic vocalization analysis in neonates and analyzed the data. R.D.P. performed part of the Fos and behavior experiments. O.I. performed the fiber photometry experiment. M.O.D. supervised the work, designed the studies, analyzed the data, and wrote the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.
REFERENCES


### KEY RESOURCES TABLE

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CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for reagents should be directed to and will be fulfilled by the Lead Contact, Marcelo Dietrich (marcelo.dietrich@yale.edu).

EXPERIMENTAL MODELS AND SUBJECT DETAILS

All preweaning mice used in the experiments were 10-21 days old from both genders. Dams were 2-6 months old. In this study, we used the following mouse lines from The Jackson Laboratories: Agrptm1(cre)Lowl/J (Agrp Cre) (JAX: 012899); B6;129P2-Gt(ROSA)26Sortm1(Trpv1,ECFP)Mde/J (R26 LSL-Trpv1) (JAX: 008513); B6.129X1-Trpv1tm1Jul/J (Trpv1 KO) (JAX: 003770); B6N.129-Rpl22tm11Psam/J (Rpl22 LSL-HA) (JAX: 011029); 129S-Npytm1Rpa/J (Npy KO) (JAX: 004545); and Slc32a1tm1Lowl (or VgatFlox/Flox) (JAX: 012897). Agrp Trpv1 mice were: AgrpCreTm/+::Trpv1—/—::R26-LSL-Trpv1Gt/+; control animals were Trpv1—/—::R26-LSL-Trpv1Gt/+ mice injected with capsaicin. Agrp Cre and R26 LSL-Trpv1 mice were backcrossed to Trpv1 KO mice to avoid the peripheral actions of capsaicin when injected systemically to activate Agrp neurons (Arenkiel et al., 2008; Dietrich et al., 2015; Güler et al., 2012; Ruan et al., 2014). We have thoroughly characterized Agrp Trpv1 mice previously (Dietrich et al., 2015). Agrp HA mice were generated by crossing Agrp Cre to Rpl22 LSL-HA mice. Analysis of ectopic expression of Cre was performed by using a specific set of primers against the excised conditional allele, as characterized before (Dietrich et al., 2015); mice with ectopic expression of the excised allele were not used in the studies. Agrp Vgat-KO mice were generated by crossing Agrp Cre to Vgat Flox/Flox mice to finally generate Agrp Cre/+:: Vgat Flox/Flox. Controls were Cre negative littermates. All mice were kept in temperature- and humidity-controlled rooms, in a 12/12 hr light/dark cycle, with lights on from 7:00 AM–7:00 PM. Food (Teklad 2018S, Envigo) and water were provided ad libitum unless otherwise stated. All procedures were approved by IACUC (Yale University).
METHOD DETAILS

Drugs
capsaicin (10 mg/kg, s.c. or i.p.; 3.33% Tween-80 in PBS; from Sigma) and oxytocin (5mg/kg, 15 IU/mg dissolved in PBS; from Sigma).

Immunohistochemistry
Mice were deeply anesthetized and perfused with freshly prepared fixative (paraformaldehyde 4%, in PBS 1x [pH = 7.4]). Brains were post-fixed overnight in fixative and sectioned on a vibratome. Coronal brain sections (50 μm) were washed several times in PBS 1x (pH = 7.4) and pre-incubated with Triton X-100 (0.3% in PBS 1x) for 30 min. Sections were then incubated in a blocking solution (Triton 0.3%, Donkey Serum 10%, Glycine 0.3M in PBS 1x) for one hour. Sections were then incubated with rabbit polyclonal anti-Fos (1:1000; #2250; Cell Signaling Technology) and mouse polyclonal anti-HA (1:1000; 901503, Biolegend) for 16 hr. After, sections were extensively washed in 0.3% Triton in PBS and incubated with secondary fluorescent Alexa antibodies (1:500). Sections were mounted and visualized by a Leica TCS SP5 Spectral Confocal Microscope (Center for Cellular e Molecular Imaging, Yale University). During the entire procedure, investigators were blinded to the experimental groups. The ImageJ analysis program (version 1.51h, NIH, USA) (Girish and Vijayalakshmi, 2004; Schindelin et al., 2012) was utilized to count the number of –HA positive (AgrpHA neurons) and Fos-positive neurons manually.

Isolation from the nest (Figures 1A–1D):
At postnatal day 10 (P10), neonates (AgrpHA) were divided into three conditions: (1) kept with the biological mother and littermates (nest); (2) isolated for 90 minutes and (3) isolated for 8 hours. Isolated animals were single-housed and placed in a clean chamber with fresh bedding. Pups were sacrificed and expression of Fos was evaluated. Samples were prepared for immunohistochemistry as described above. AgrpHA pups were used in these studies to allow identification of Agrp neurons. All samples were prepared and counted blinded for the experimental groups.

Milk intake in the nursing nest (Figures 1E and 1F):
when ten days of age (P10), pups were divided into two groups: kept with the biological mother or isolated for 90 minutes. Isolated pups were reintroduced to the biological mother for another 90 minutes after isolation. Body weight was measured prior and after 90 minutes. To ensure that animals within each group had similar milk availability, they were tested using an equal number of animals. Upon reintroduction of the isolated pups to the home cage, pups that stayed with the dam were removed to avoid competition. These experiments were not blinded.

Measurement of corticosterone levels (Figure 1G):
At postnatal day 10, neonates were divided into two groups: kept with the biological mother or isolated for 90 minutes. After testing, neonates were deeply anesthetized, and blood samples were collected through cardiac puncture. A total of 150 μL of blood was collected. Blood samples were left at room temperate for one hour and centrifuged at 5000 rpm for 20 minutes. Plasma was collected and stored at −80°C. Corticosterone level was measured using an enzyme immunoassay (ELISA) kit (Enzo Life Sciences, Farmingdale, NY, USA) according to the manufacturer’s instructions.

Isolation from the nest in the presence of a predator odor (Figures 1H–1J):
At postnatal day 10 (P10), pups (AgrpHA) were divided into three conditions: (1) kept with the biological mother and littermates (nest); (2) isolated for 90 minutes and (3) isolated for 90 minutes in the presence of a synthetic predator odor (2,4,5-Trimethylthiazole (mT), Sigma-Aldrich). Isolated animals were single-housed and placed in a clean chamber with fresh bedding. Ten microliters of mT odor were pipetted onto a small square nesting material (2 × 2 cm). To avoid contact with the odor, the chamber was divided by a wire mesh resulting in two small compartments, allowing the pups to smell the odor.

Isolation from the nest with milk infusion (Figures 1K–1O):
The procedure of milk infusion consisted in the insertion of a polyurethane-based catheter tubing (Micro-Renathane® Tubing, MRE-033, Braintree Scientific, Inc.) attached to a pump. To avoid an invasive procedure, the inserted tube end was heated and bent to create a small U shape at its end tip. After insertion into the mouth, the tube was attached to the fur on the outside of the cheek using a small drop of crazy glue to hold it in place. The whole procedure did not require anesthesia and last less than 30 s. A total of 200 μL of milk was infused during the 90 minutes (15 μL ejections, every 5-15 minutes). The following types of milk were used: (1) commercial Half & Half cow’s milk (Organic Valley, Ultra Pasteurized Grade A); an (2) mouse milk collected from lactating dams. To confirm that milk was infused, a tasteless blue dye (Erioglaucine disodium salt, Sigma-Aldrich, Cat. 861146) was added (< 1 mg/mL) to the milk prior infusion. The stomach was excised to confirm the blue color indicating milk ingestion.
Mouse milk collection

The milk collection was performed on lactating dams with litters between the ages of P8-P12. Dams were separated from their litter for 6 hours prior to collection to ensure adequate milk production. Dams were lightly anesthetized with isoflurane and oxytocin (5 mg/kg, i.p) was administered to promote milk release. Milk was expressed from the nipples using pressure from the thumb and forefinger to gently massage and squeeze the mammary tissue in an upward motion until a visible bead of milk begins to form at the base of the teat. Then, milk was collected using a 20 μL calibrated pipette, pipetted into a 1.5 mL Eppendorf tube and stored at −20°C until the day of the test. The duration of the milk collection lasts less than 10 minutes.

Artificial feeding protocol (Figures 1P–1R):

when seven days old, neonates were separated from the biological dam and kept with a non-lactating foster dam. Every 3-4 hours, the neonates were separated from the foster dam and milk was provided for ~30 minutes by the experimenter using a surrogate nipple attached to a tip in a 100 μL pipette. The volume of intake in each session varied between 80 μL to 150 μL of milk. Because of the limitation in getting mouse milk, we performed the artificial feeding using a more caloric formula of cow milk (Heavy Whipping Cream, Organic Valley) that resembles the nutrition facts of a mouse milk (Gors et al., 2009). At postnatal day 10, milk was provided for 30 minutes and immediately after neonates were separated into two groups: kept with the non-lactating foster dam and littermates (nest) or isolated for 90 minutes. All other procedures for Fos counting were as described above.

Assessment of maternal components with foster dams (Figures 2A–2C):

At postnatal day 10, neonates were separated from the biological dam and placed in the cage of a foster dam. Foster dams in different lactation conditions were used: (1) non-lactating foster dam; (2) non-lactating foster dam with protruded nipples; (3) lactating foster dam. Dam rodents have the nipples still distended without milk release permitting suckling for two weeks after weaning if the female is not pregnant again. Lactating foster dams were chosen in a similar postnatal day of lactation, and their offspring was removed immediately before placing the alien/unfamiliar neonates. Neonates were divided into five groups: (1) kept with the biological dam and littermates (nest); (2) kept with a non-lactating foster dam; (3) kept with a non-lactating foster dam with protruded nipple; (4) kept with lactating foster dam with protruded nipple and (5) isolated for 90 minutes. All other procedures for Fos counting were as described above.

Isolation from the nest with thermal support (Figures 2D–2G):

When ten-day-old, neonates were separated from the biological dam, and thermal support was provided using two different conditions. In the first condition, neonates were placed in a humidity and temperature-controlled climate chamber (70%–80% of humidity, 35°C, Sables Systems). In the second condition, a thermal support device set at 35°C was placed underneath the chamber in which the neonates were separated. We confirmed appropriated thermal conditions by monitoring the temperature throughout testing using calibrated thermometers. Neonates were divided into four groups: (1) kept with the biological dam and littermates (nest); (2) isolated for 90 minutes at thermoneutrality; (3) isolated for 90 minutes in the thermal support device; and (4) isolated for 90 minutes without thermal support (room temperature). All other procedures for Fos counting were as described above.

Isolation from the nest in pups raised at thermoneutrality (Figures 2H–2J):

The lactating dam was placed in a humidity and temperature-controlled climate chamber (70%–80% of humidity, 35°C, Sables Systems) two weeks before delivery to acclimate to the new environment. Temperature and humidity in the climate chamber were monitored twice a day until testing. At postnatal day 10, neonates were divided into three groups: (1) kept with the biological dam and littermates (nest); (2) isolated for 90 minutes at room temperature; and (3) isolated for 90 minutes at thermoneutrality. All other procedures for Fos counting were as described above.

Fiber photometry (Figure 3):

Agp2Cre/Cre mouse neonates (P0–P1) were cryo-anesthetized. Neonates were placed on ice, using aluminum foil as a barrier to prevent direct contact with the ice. After 8 minutes, neonates were removed from the ice and placed onto a chilled rat/mouse neonatal frame (Stoelting Co., Wood Dale, IL). A Cre-dependent adeno-associated virus (AAV) encoding the calcium sensor GCaMP7s (AAV8-CAG-Flex-GCaMP7s-SV40, Penn Vector Core) was injected unilaterally at a volume of 300 nL using following coordinates from lambda: AP = +.98 ML, lateral = −.3mm, DV = −.41mm. On postnatal day 12, a fiber optic cannula (NA = 0.48, core diameter = 400 μm from Doric Lenses) was placed over the arcuate nucleus using following coordinates from bregma: AP = −1.38 mm, lateral = −0.3mm, DV = −5.8 mm. One to two days after placing the fiber optic cannula, experimental mice were placed in a Plexiglas cage (10 cm x 8 cm x 6 cm) with 4 siblings and bedding from home cage. After 5 minutes of baseline fiber photometry recordings (see below), mice connected to the fiber photometry system were moved to an identical adjacent Plexiglas cage for a period of ten minutes of isolation. Subsequently, experimental mice were return to the cage with the siblings for 5 minutes. The fiber photometry system consisted of two different sets of LEDs: 405 nm LED sinusoidally modulated at 211 Hz and a 460 nm LED sinusoidally modulated at 333 Hz. Both light streams were merged into an optical fiber patch using a minicube (Doric Systems). The fiber optic patch was connected to the cannula on the mouse pup. Fluorescence emitted by jGCaMP7s in response to light excitation was collected with same fiber patch cord and focused into a photodetector (Newport). The signal collected at the photodetector was
collected in a digital fiber photometry processor (RZ5P, Tucker-Davis Technologies). Signal was processed and pre-analyzed using the Synapse Software Suite (Tucker-Davis Technologies). The data were exported to MATLAB for post-processing. First, the isosbestic channel (405 nm excitation) was fitted to the calcium-dependent channel (460 nm excitation, denoted as F) using first order polynomial fitting (F$_0$ denotes the fitted isosbestic). The calcium fluorescence activity was calculated as: (F - F$_0$)/F$_0$. The high-frequency components of the fluorescence activity were then filtered out by a low pass filter at 0.5 Hz. We then down sampled the signal by averaging it in non-overlapping windows of 0.1 s. The Z-score was calculated considering the minute before isolation as the baseline.

**Recording of ultrasonic vocalizations (Figures 4 and 5):**
P10 mice were separated from the dam and placed in a soundproof chamber. In the thermoneutrality experiment, pups were divided into two groups: (1) isolated for 90 minutes at room temperature or (2) isolated for 90 minutes at thermoneutrality (70%–80% of humidity, 35°C, Sables Systems). USVs were recorded for 90 minutes. In experiments with the Npy$^{K0}$ and Agrp$^{Trpv1-K0}$ mice, USVs were recorded for ten minutes immediately following separation from the dam in the soundproof chamber. In the experiment with Agrp$^{Trpv1}$ mice, this initial ten minutes was considered as a baseline before activation of Agrp neurons. Then, P10 mice were injected with capsaicin (10 mg/kg, s.c) and USVs were recorded for an additional twenty minutes. USVs were recorded using an UltraSoundGate Condenser Microphone CM 16 (Avisoft Bioacoustics, Berlin, Germany) placed 10 cm above the animals. The microphone was connected via an UltraSoundGate 416 USGH audio device and recorded with a sampling rate of 250,000 Hz by the software Avisoft RECORDER (version 4.2.16; Avisoft Bioacoustics).

**Maternal preference test (Figures 5N–5P):**
The maternal preference test was performed in a three-chamber apparatus (65 × 42 × 23 cm) and comprised of three stages: Stage 1 – acclimation: the dam was allowed to explore the apparatus without the presence of pups for ten minutes. Stage 2 – exploration: two P10 mice (control and Agrp$^{Trpv1}$, n = 25 pairs) were placed on each side of the apparatus inside of an inverted metal wire cup and the dam was allowed to explore the pups for ten minutes. Stage 3 - preference: dam was restricted to the center compartment, pups were injected with capsaicin (10 mg/kg, s.c) and USVs were recorded in the cups and then the dam was allowed to explore all compartments for 20 minutes. Groups were randomly alternated between both sides to avoid preference for one side of the chamber. Time spent interacting with the pups was measured using Any-maze (Stoelting Co., Wood Dale, IL).

**Analysis of ultrasonic vocalizations**
Ultrasonic vocalizations (USV) were automatically extracted from the audio recordings by using spectral analysis through image processing. Each audio file was analyzed in segments of 1 minute long and then Short Fourier transformed with a Hamming windowing function (window = 256), NFFT = 1024 sampling points and an overlap between successive windows equal to half of the window size. These parameters generate a spectrogram with resolution of 0.5 ms and 244 Hz. The spectrograms were converted to grayscale images and the USVs were segmented on the spectrogram through a sequence of image processing techniques, which included the contrast enhancement of the image ($\gamma = 1$), the application of an adaptive threshold (sensitivity of 20%) followed by a series of morphological operations and identification of connected components. The segmented USV candidates were then analyzed by a local median filtering (LMF) to eliminate segmentation noise based on the contrast between an USV candidate and its background. The minimum contrast acceptable between an USV candidate and its background was automatically estimated based on a differential geometry analysis of the contrast of all the USV candidates detected in an audio recording. USVs less than 10 ms apart were considered as part of the same syllable. Next, all the USVs were classified in 11 distinct call types (Grimsley et al., 2011) by a Convolutional Neural Network, which had the AlexNet architecture as starting point. The network was trained for USV classification with over 14,000 samples of real USVs, which were then augmented in order to increase the variability of the samples, resulting in > 57,000 samples. The output consisted of a table summarizing the main features of the USVs detected. This table contains the start and end time of the USVs, as well as its mean, maximum and minimum frequency, mean intensity and other relevant spectral features such as the existence of harmonic components. Each vocalization received a label based on the most likely call type label attributed by the Convolutional Neural Network. The label of each USV is also available as a probability distribution function over all the call types. The software was custom developed in our laboratory and is available upon request. The details of the software will be published elsewhere.

**Mouse pup behavior toward the anesthetized dam (Figures 6A–6G and 7A–7J):**
Animals were recorded under infrared illumination and assessed for 20 minutes at postnatal day 10 (P10) and postnatal day 15 (P15). Each animal was tested at one age only. Before the experiment, the dam was anesthetized (100 mg/kg Ketamine + 10 mg/kg Xylazine). The maximum number of pups tested per dam was eight. Animals received an injection of capsaicin (10 mg/kg, s.c.) before the experiment. We used a custom built-chamber (20 × 15 cm built in opaque Plexiglas). The dam was placed at an angle of 45° on her back along the edge. Pups were placed on the other edge of the chamber, ~20 cm away from their dam. Parameters such as latency to attach to the dam’s nipple, distance traveled, and the number of nipple attachments were assessed using Any-Maze (Stoelting Co., Wood Dale, IL). Experiments were performed blinded for the genotype.
Mouse pup behavior toward the anesthetized dams injected with oxytocin (Figures 6P–6V and 7K–7P):

In anesthetized dams, milk ejection is largely decreased, and dams are considered non-lactating (Lincoln et al., 1973). To circumvent this issue, we performed a similar experiment as described above but injected the anesthetized dams with oxytocin (5 mg/kg, i.p) immediately before each test. Nipples were manually expressed to confirm that there was milk ejection before the experiment. P10 mice received an injection of capsaicin (10 mg/kg, s.c) and were subsequently assessed for 20 minutes. In this assay, a second injection of oxytocin (5 mg/kg, i.p) was given at 10 minutes of test. P15 mice received an injection of capsaicin (10 mg/kg, s.c) and were subsequently assessed for 10 minutes. The duration of the testing period was shorter because P15 mice quickly attached to the dam’s nipple in preliminary experiments. Parameters such as latency to attach to the dam’s nipple, distance traveled, and the number of nipple attachments were analyzed using Any-Maze (Stoelting Co., Wood Dale, IL). Body weight was measured prior and after testing. Experiments were performed blinded for the genotype.

Independent feeding (Figures 7Q and 7R):

Mice were tested at postnatal days P15, P18, and P21. Naive animals were used for each postnatal age and mice were acclimated to the behavior room for one-hour before the experiment. Food was left inside the cage to prevent a state of deprivation. Animals were tested in a mouse cage filled with home bedding and two Petri dishes placed in opposite corners. After the acclimation period (1 hour), the experiment was performed. Animals were removed from the cage, received an injection of capsaicin (10 mg/kg, i.p.) and were returned to the cage. One Petri dish was empty; the other had a pellet of chow diet. Body weight and food intake were evaluated after 30 minutes. Experiments were performed blinded for the genotype.

QUANTIFICATION AND STATISTICAL ANALYSIS

MATLAB (2016a or above) and Prism 8.0 were used to analyze data and plot figures. All figures were edited in Adobe Illustrator CS6/CC. Illustrations were designed by Mind the Graph (MindtheGraph.com). Data were first subjected to a normality test using the D’Agostino & Pearson normality test or the Shapiro-Wilk normality test. When homogeneity was assumed, a parametric analysis of variance test was used. The Student’s t test was used to compare two groups. Welch’s correction was used when standard deviations were unequal between groups. ANOVA was used to compare multiple groups. Tukey-Kramer’s multiple comparisons test was used to find post hoc differences among groups and calculate 95% confidence intervals to report effect size. When 95% confidence intervals were not calculated, then the Holm-Sidak’s multiple comparisons test was used. When homogeneity was not assumed, the Kruskal-Wallis nonparametric ANOVA was selected for multiple statistical comparisons. The Mann-Whitney U test was used to determine significance between groups. Two sample Kolmogorov–Smirnov test was used to calculate the statistical differences between features of ultrasonic vocalizations. Chi-square test was used to find differences in the number of pups that attached to nipples in the behavior tests performed in neonates. One- or two-tail tests were used based on prior experimental hypothesis. Statistical data are provided in text and in the figures. In the text, values are provided as mean ± SEM p < 0.05 was considered statistically significant.
Figure S1. PCR-Based Analysis for Genotyping Agrp<sup>Trpv1</sup> and Control Mice, Related to Figure 5

(A) Illustrative diagram of the tissue collection and PCR-based analysis genotyping.

(B) Genomic DNA samples from P15 mice were extracted and amplified using primers for Trpv1 knockout allele. The lower band is the knockout allele for the Trpv1 gene.

(C) Genomic DNA samples from P15 mice were extracted and amplified using primers for ectopic Trpv1 allele. The upper band (667 bp) shows that the excised allele for Trpv1 is specifically expressed in the arcuate nucleus of the hypothalamus.