

1 **Rapid shift in substrate utilization driven by hypothalamic Agrp neurons**

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1 **Abstract**

2 Agrp neurons drive feeding. To what extent these neurons participate in the regulation of other
3 homeostatic processes is not well understood. We investigated the role of Agrp neurons in
4 substrate utilization in mice. Activation of Agrp neurons was sufficient to rapidly increase RER
5 and carbohydrate utilization, while decreasing fat utilization. These metabolic changes were
6 linearly correlated with carbohydrates ingested, but not protein or fat ingestion. However, even
7 in the absence of ingestive behaviors, activation of Agrp neurons led to changes in substrate
8 utilization in well-fed mice. These effects were coupled to metabolic shifts towards lipogenesis.
9 Inhibition of fatty acid synthetase (FAS) blunted the effects of Agrp neurons on substrate
10 utilization. Finally, Agrp neurons controlled peripheral metabolism, but not food intake, via β 3-
11 adrenergic receptor signaling in fat tissues. These results reveal a novel component of Agrp
12 neuron-mediate metabolism regulation that involves sympathetic activity on fat compartments to
13 shift metabolism towards lipogenesis.

1 **Introduction**

2 Obesity is a major health problem that results from altered regulation of energy
3 metabolism. The central nervous system tightly controls energy metabolism by regulating
4 hormonal and autonomic action on peripheral tissue. The hypothalamus is a conserved region in
5 the brain involved in homeostatic control, including energy balance. The arcuate nucleus of the
6 hypothalamus contains a population of neurons that selectively expresses agouti-related protein
7 (AgRP; hereafter, Agrp neurons) ([Broberger et al., 1998](#); [Hahn et al., 1998](#); [Ollmann et al., 1997](#);
8 [Rossi et al., 1998](#)). Agrp neurons are located in proximity to the third ventricle and have direct
9 access to circulation ([Olofsson et al., 2013](#)), as this brain region lacks a complete blood-brain
10 barrier ([Broadwell et al., 1983](#); [Broadwell and Brightman, 1976](#)). Consistently, Agrp neurons are
11 known to respond to a variety of circulating factors ([Könner et al., 2007](#); [Pinto et al., 2004](#);
12 [Steculorum et al., 2015](#); [van den Top et al., 2004](#)). However, recent evidence demonstrates that
13 Agrp neurons also integrate food-related information via sensory pathways ([Betley et al., 2015](#);
14 [Chen et al., 2015](#); [Mandelblat-Cerf et al., 2015](#)). Thus, Agrp neurons are unique as they gather
15 relevant information from several sources to regulate physiology and behavior.

16 Agrp neurons were first described to be active during food deprivation, a phenomenon
17 conserved across rodent and non-human primate species ([Grove et al., 2003](#); [Hahn et al., 1998](#);
18 [Mandelblat-Cerf et al., 2015](#); [Takahashi and Cone, 2005](#)). Because Agrp neurons co-express
19 NPY ([Broberger et al., 1998](#); [Hahn et al., 1998](#)), and because NPY acts as a potent orexigenic
20 peptide when injected into the brain ([Clark et al., 1984](#)), it was natural to conclude that during
21 food deprivation Agrp neurons are active to drive food intake. Subsequent work demonstrated
22 that activation of Agrp neurons is sufficient to drive food intake in sated mice ([Aponte et al.,](#)

1 [2011](#); [Dietrich et al., 2015](#); [Krashes et al., 2011](#)). Conversely, elimination of Agrp neurons in the
2 adult brain led to aphagia ([Gropp et al., 2005](#); [Luquet et al., 2005](#)).

3 Recent work demonstrated a specific role for Agrp neurons in metabolic processes,
4 including the control of white adipose tissue (WAT) browning and thermogenesis ([Ruan et al.,](#)
5 [2014](#)), as well as brown adipose tissue (BAT) glucose metabolism ([Steculorum et al., 2016](#)).

6 Here, we investigate the role of Agrp neurons in the control of peripheral substrate utilization.

7 Our data demonstrate that Agrp neuron activation rapidly regulates peripheral metabolism

8 independently of food ingestion. Specifically, Agrp neuron activation promotes lipogenesis via

9 sympathetic nervous system (SNS). These results highlight the complex function of these key

10 hypothalamic neurons in normal physiology and in disordered metabolic states, such as obesity.

1 **Results**

2 *Acute switch in nutrient utilization upon Agrp activation*

3 Nutrient utilization can be measured by indirect calorimetry, where the measurements of
4 VCO₂ production and VO₂ consumption are used to calculate the respiratory exchange ratio
5 (RER) (**Figure 1A**) (Frayn, 1983). Under normal conditions, a RER approaching 0.7 indicates
6 predominant fat oxidation, while a RER approaching 1.0 indicates predominant carbohydrate
7 oxidation (**Figure 1A**) (Frayn, 1983). To test for the acute role of Agrp neurons in nutrient
8 utilization we took advantage of an animal model that we have recently characterized (Dietrich et
9 al., 2015; Ruan et al., 2014), in which Agrp neurons are transiently activated by peripheral
10 injection of capsaicin in *Agrp*^{Trpv1} mice (Dietrich et al., 2015; Ruan et al., 2014). *Agrp*^{Trpv1} mice
11 are generated by crossing *Agrp*^{Cre} to *Rosa26*^{LSL-Trpv1} mice backcrossed to a *Trpv1*^{KO} background
12 to prevent capsaicin action on other cell types (Arenkiel et al., 2008; Guler et al., 2012). As a
13 result, *Agrp*^{Trpv1} mice selectively express the capsaicin-sensitive channel, Trpv1, in Agrp
14 neurons. Because capsaicin is a highly specific ligand of Trpv1 (Caterina et al., 1997), peripheral
15 injection of capsaicin allows for a rapid, reliable and transient chemogenetic activation of Agrp
16 neurons in *Agrp*^{Trpv1} mice (Dietrich et al., 2015; Ruan et al., 2014). Using *Agrp*^{Trpv1} mice allowed
17 us to rapidly activate Agrp neurons without the necessity of tethers (as for example, using
18 optogenetics), facilitating the study of animals in indirect calorimetry chambers to measure gas
19 (O₂ and CO₂) exchange. Importantly, Trpv1-mediated activation of Agrp neurons is both rapid
20 (latency to start is ~2 min) and short lived (lasts ~1h), unlike the effects of activating Agrp
21 neurons via the designer receptor hM3Dq (Krashes et al., 2011).

22 We injected mice with capsaicin during the light cycle and recorded concomitant changes
23 in locomotor activity and gas exchange for the ensuing 24 hours. Capsaicin injections produced

1 sharp increases in RER in *Agrp*^{Trpv1} mice, (**Figure 1B**), in line with the previously observed fast
2 kinetics of feeding induced by *Agrp* neuronal activation in this animal model ([Dietrich et al.,](#)
3 [2015](#)). We did not observe statistically significant changes in either VO₂ (**Figure 1C**), VCO₂
4 (**Figure 1D**) or energy expenditure (**Figure 1E**). Because exercise increases RER, we also
5 measured activity levels upon *Agrp* neuron activation. In line with our previous report ([Dietrich](#)
6 [et al., 2015](#)), activation of *Agrp* neurons in the home cage in the presence of food did not
7 increase levels of ambulatory activity (**Figure 1F**), indicating that the increase in RER was not
8 affected by elevated locomotion. Based on gaseous exchange ([Frayn, 1983](#)), we calculated total
9 rates of fat utilization and carbohydrate utilization for the whole animal. In line with changes in
10 RER, activation of *Agrp* neurons led to a rapid and prolonged decrease in fat utilization (**Figure**
11 **1G**) concomitant with an increase in carbohydrate utilization (**Figure 1H**). Because (i) this
12 experiment was performed in the presence of food and (ii) the effects on nutrient utilization were
13 prolonged compared to feeding upon *Agrp* neuron activation ([Dietrich et al., 2015](#)), we
14 hypothesized that these metabolic shifts are due to changes in postprandial metabolism.

15 *Effects of diet ingestion on metabolic switches*

16 To further dissect the effects of altered RER from increased food intake upon *Agrp*
17 neuron activation, we determined how diet composition affects the metabolic phenotypes by
18 varying the proportion of macronutrients in the diet (**Figure 2A**; see Material and Methods for
19 more details). We activated *Agrp* neurons in *Agrp*^{Trpv1} mice during the light cycle, providing
20 controlled amounts of different diets: LFHS, a low-fat high-sugar diet containing 10% kcal from
21 fat sources; HF45, a high-fat diet containing 45% kcal from fat sources; and HF60, a high-fat
22 diet containing 60% kcal from fat sources. The three diets contain 20% kcal from protein
23 sources. We performed these experiments in three different conditions: (i) limiting the amount of

1 calories ingested from fat sources (0.4 kcal from fat, ‘Fat Clamp’); (ii) limiting the amount of
2 calories ingested from carbohydrate sources (0.7 kcal from sugars, ‘Sugar Clamp’), or (iii)
3 feeding animals isocaloric quantities of food (1.5 kcal from all sources, ‘Cal Clamp’).

4 Before activation of *Agrp* neurons, baseline RER was 0.79 ± 0.003 (mean \pm s.e.m.; $n =$
5 90 measurements), fat utilization was $0.51 \pm 0.01 \text{ mg} \cdot \text{min}^{-1}$ (mean \pm s.e.m.; $n = 90$
6 measurements) and carbohydrate utilization was $0.61 \pm 0.02 \text{ mg} \cdot \text{min}^{-1}$ (mean \pm s.e.m.; $n = 90$
7 measurements), indicating that in the conditions tested animals were using substrates from mixed
8 sources. Following activation of *Agrp* neurons, we observed an increase in RER in all conditions
9 tested (**Figure 2B** and **Figure 2 - Figure supplement 1A**). In linear regression analyses, we
10 found a strong correlation between RER and carbohydrate ingestion ($r^2 = 0.85$, $P < 0.001$), but
11 not protein ($r^2 = 0.30$, $P = 0.12$) or fat ($r^2 = 0.12$, $P = 0.35$) (**Figure 2C**). Fat utilization was
12 reduced upon *Agrp* neuron activation in all conditions tested (**Figure 2D** and **Figure 2 – Figure**
13 **supplement 1F**) and was negatively correlated with carbohydrate ingestion ($r^2 = 0.85$, $P <$
14 0.001), but not protein ($r^2 = 0.26$, $P = 0.15$) or fat ($r^2 = 0.16$, $P = 0.25$) (**Figure 2E**).
15 Carbohydrate utilization followed changes in RER, increasing upon *Agrp* neuron activation in all
16 dietary conditions tested (**Figure 2F** and **Figure 2 - Figure supplement 1G**). Changes in
17 carbohydrate utilization were strongly correlated to carbohydrate ingestion ($r^2 = 0.90$, $P <$
18 0.0001 ; **Figure 2G**). Protein ingestion also correlated with carbohydrate utilization, but linear
19 regression analysis did not reach statistical significance ($r^2 = 0.39$, $P = 0.07$; **Figure 2G**). Fat
20 intake did not correlate with carbohydrate utilization ($r^2 = 0.07$, $P = 0.46$; **Figure 2G**). As
21 expected, these results demonstrated that carbohydrate levels in the diet are main players in the
22 metabolic shift that occurs upon food ingestion, suggesting that macronutrient quality rather than
23 caloric content account for these effects.

1 *Agrp* neurons control substrate utilization in the absence of ingestion

2 The effect of *Agrp* neuron activation on carbohydrate metabolism led us to further
3 explore the extent to which activation of *Agrp* neurons and sugar ingestion interact to rapidly
4 shift metabolism. We provided different concentrations of glucose to mice via gavage delivery,
5 thereby ruling out potential cephalic phase effects (**Figure 3A**). Gavage infusion of glucose led
6 to dose-dependent increases in RER (Figure 3B; $r^2 = 0.99$, $P = 0.003$), decreases in fat utilization
7 (**Figure 3C**; $r^2 = 0.99$, $P = 0.004$), and increases in carbohydrate utilization (**Figure 3D**; $r^2 =$
8 0.99 , $P = 0.003$). These metabolic shifts were mainly due to increases in VCO_2 (**Figure 3F**; $r^2 =$
9 0.98 , $P = 0.006$) and not VO_2 (**Figure 3E**; $r^2 = 0.84$, $P = 0.07$). Glucose intake also positively
10 correlated with energy expenditure (**Figure 3G**; $r^2 = 0.92$, $P = 0.03$), likely due to the thermic
11 effects of carbohydrate digestion. Activity levels were unchanged (**Figure 3D**; $r^2 = 0.58$, $P =$
12 0.23). These experiments demonstrate that small amounts of carbohydrate ingestion alone are
13 sufficient to shift metabolism even in the absence of carbohydrate sensing at the level of the
14 mouth. Thus, our previous results on substrate metabolism upon activation of *Agrp* neurons
15 could be simply explained by ingestion of carbohydrates. We designed the next experiments to
16 test this consideration.

17 We infused both control and *Agrp*^{Trpv1} mice with glucose (2 g/kg body weight, via
18 gavage) and activated *Agrp* neurons by injecting capsaicin (**Figure 4A**). Strikingly, activation of
19 *Agrp* neurons led to an increased peak and a more sustained elevation of RER (**Figure 4B**), an
20 effect that was present even in mice infused with saline (**Figure 4B**). Accordingly, activation of
21 *Agrp* neurons led to a sustained decrease in fat utilization (**Figure 4C**) and increase in
22 carbohydrate utilization (**Figure 4D**), regardless of glucose infusion. These results strongly
23 suggest that activation of *Agrp* neurons alone is sufficient to promote shifts in substrate

1 utilization, independently of food consumption. However, even in animals infused with saline,
2 small amounts of liquid were delivered via gavage, raising the possibility that gastric distension
3 acts together with Agrp neuron activation to promote changes in substrate utilization. To exclude
4 this possibility, we repeated our experiments in a new cohort of mice in which Agrp neurons
5 were activated in the absence of food (**Figure 4 supplement 1A**). In line with our previous
6 findings, activation of Agrp neurons alone was sufficient to increase RER (**Figure 4 supplement**
7 **1B**), while decreasing fat utilization (**Figure 4 supplement 1G**) and increasing carbohydrate
8 utilization (**Figure 4 supplement 1H**). The above experiments demonstrate that Agrp neurons
9 rapidly control whole body substrate utilization by shifting metabolism towards carbohydrate
10 relative to fat utilization independently of ingestive behaviors.

11 *Participation of lipogenesis in Agrp neuron-mediated shifts in metabolism*

12 A physiological scenario in which the metabolic shift promoted by Agrp neuron
13 activation is expected to be important is during positive energy balance (i.e., a metabolic state
14 coupled to weight gain), as favoring carbohydrate utilization would allow storage of lipids and
15 re-route of energy substrates to undergo de novo lipogenesis, further increasing fat deposition. In
16 line with this, we found that, in absence of food ingestion, Agrp neuron activation decreased
17 circulating levels of non-esterified fatty acids (NEFAs, **Figure 5B**) with no changes in blood
18 glucose levels (**Figure 5C**) in well-fed mice (**Figure 5A**). Because circulating NEFAs decrease
19 upon Agrp neuron activation, these results suggest a decrease in release, and possibly an increase
20 in deposition. To test this hypothesis, we measured expression levels of genes involved in lipid
21 metabolism in the white adipose tissue (WAT) from *Agrp*^{Trpv1} and control mice 60 minutes after
22 capsaicin injection. We found a decrease in the expression level of *Ppara* (**Figure 5D**), a gene
23 involved in the promotion of fat catabolism. We also found a significant increase in expression

1 levels of *hexokinase2* (*hk2*; **Figure 5D**), a rate-limiting enzyme involved in glycolysis, a critical
2 metabolic step to provide carbons for de novo lipogenesis. Hormone-sensitive lipase (HSL) is an
3 essential step in the breakdown of triglycerides to release fatty acids in circulation. Activation of
4 HSL occurs by phosphorylation of this enzyme in several serine residues. Upon activation of
5 *Agrp* neurons, we found decreased levels of phosphorylated HSL (pHSL) in WAT compartments
6 (**Figure 5E**). Together, these experiments indicate activation of *Agrp* neurons leads to increased
7 lipogenesis and decreased lipolysis in the WAT.

8 De novo lipogenesis can drive RER above 1.0 (Frayn, 1983), and could be a potential
9 factor involved in *Agrp* neuron mediated acute shifts in RER. Additionally, in our dietary clamp
10 experiments (**Figure 2**), we found negative results for fat utilization in mice that ate large
11 amounts of sugars (**Figure 2 supplement 1F**). Because calculated fat utilization is the sum of
12 true rates of fat oxidation minus the rate of synthesis of fat from carbohydrates (de novo
13 lipogenesis) (Frayn, 1983), an increase in synthesis leads to a net decrease in calculated fat
14 utilization. Thus, negative values for calculated fat utilization *only* occur when the rate of
15 synthesis is higher than the rate of oxidation (Frayn, 1983), and are pathognomonic of ongoing
16 lipid synthesis. To test for the participation of fat synthesis in the rapid effects of *Agrp* neurons
17 on metabolism, we blocked fatty acid synthetase (FAS), a key enzyme involved in fat storage
18 (Lodhi et al., 2012). We treated mice with a pharmacological inhibitor of FAS (C75, 10 mg/kg,
19 i.p.) (Kuhajda et al., 2000; Loftus et al., 2000) and activated *Agrp* neurons in indirect calorimetry
20 chambers (**Figure 5F**). Treatment of control mice with the FAS inhibitor had no effects on RER
21 (**Figure 5G**), fat utilization (**Figure 5H**) or carbohydrate utilization (**Figure 5I**). The lack of
22 effects of FAS inhibition on substrate utilization is in line with the low levels of lipogenesis
23 during the light cycle of mice. In contrast to control animals, inhibition of FAS blocked the

1 effects of *Agrp* neuron activation on substrate utilization (**Figures 5G-I**). These results
2 demonstrate *Agrp* neurons rapidly shift metabolism towards lipogenesis in well-fed animals.

3 *Sympathetic signaling mediates peripheral effects of Agrp neurons*

4 We next determined if *Agrp* neurons control peripheral substrate utilization via SNS
5 signaling. Norepinephrine release and binding to adrenergic receptors on fat compartments
6 promotes lipolysis, while its inhibition favors lipogenesis. Accordingly, we predicted that *Agrp*
7 neurons control adiposity by inhibiting sympathetic signaling on fat compartments and
8 promoting lipogenesis in anabolic states. To test for this hypothesis, we treated mice with a β 3-
9 adrenergic receptor agonist (CL 316,243) ([Bloom et al., 1992](#)) (**Figure 6A**), as β 3-adrenergic
10 receptors are highly selective to fat compartments ([Grujic et al., 1997](#)). Treatment of control
11 mice with CL 316,243 did not alter RER (**Figure 6B**), but prevented the increase in RER upon
12 *Agrp* neuron activation in *Agrp*^{Trpv1} mice (**Figure 6B**). When we calculated fat utilization, CL
13 316,243 completely reverted the inhibition of whole body fat utilization upon *Agrp* neuron
14 activation in *Agrp*^{Trpv1} mice but had no effects in control animals (**Figure 6C**). Concomitantly,
15 CL 316,243 prevented the increase in carbohydrate utilization upon activation of *Agrp* neurons
16 (**Figure 6D**). These results demonstrate that promotion of β 3-adrenergic receptor signaling in fat
17 compartments is sufficient to revert the effects of *Agrp* neurons on peripheral fuel metabolism.

18 Because *Agrp* neurons are characteristically linked to food intake control ([Aponte et al.,](#)
19 [2011](#); [Gropp et al., 2005](#); [Hahn et al., 1998](#); [Krashes et al., 2011](#); [Luquet et al., 2005](#); [Rossi et al.,](#)
20 [1998](#)), we investigated whether the effects of *Agrp* neurons on fuel utilization are
21 mechanistically linked to feeding. Thus, we tested whether CL 316,243 could acutely block the
22 effects of *Agrp* neuron activation on food intake. We first performed a dose response study to
23 detect the range of CL 316,243 doses that could revert the increase in RER upon *Agrp* neuron

1 activation. We found that doses as low as 0.01 mg/kg almost completely reverted the effects of
2 Agrp neurons on RER (**Figure 6E**). Next, we selected two doses of CL 316,243 (1.00 and 0.01
3 mg/kg) to investigate its effects on Agrp neuron mediated food intake. CL 316,243 is highly
4 anorexigenic ([Grujic et al., 1997](#)), but its effects are observed after several hours and not as rapid
5 as the effects of Agrp neurons on feeding ([Aponte et al., 2011](#); [Dietrich et al., 2015](#); [Krashes et](#)
6 [al., 2011](#)). In all conditions tested, CL 316,243 did not revert the effects of Agrp neuron
7 activation on food intake (**Figures 6F-G**). These latter findings indicate a divergence between
8 the feeding and metabolic mechanisms underlying Agrp neuron function and support the
9 argument that Agrp neurons favor the storage of fat (lipogenesis) in situations of energy surfeit
10 (**Figure 6H**).

1 **Discussion**

2 We reported here a mechanism by which elevated Agrp neuron activity shifts metabolism
3 towards lipid storage via the SNS. The SNS releases norepinephrine in target organs to regulate a
4 variety of physiological functions. In WAT, which stores excess of energy in the form of fat,
5 activation of the SNS leads to lipolysis ([Bartness et al., 2014](#); [Correll, 1963](#); [Zeng et al., 2015](#)),
6 breaking down triglycerides to release free fatty acids in the circulation ([Zeng et al., 2015](#)).
7 Lipolysis is critical for survival during periods of food scarcity, with fatty acids becoming the
8 predominant energy substrate. Conversely, inhibition of lipolysis favors lipogenesis, leading to
9 fat deposition ([Rutkowski et al., 2015](#)). This is an important adaptive response to allow storage
10 of the excess of energy for later mobilization during food deprivation. However, when this
11 anabolic state (lipogenesis > lipolysis) is sustained it can become maladaptive and trigger
12 obesity. Here, we provided evidence that elevated activity of Agrp neurons is sufficient to
13 promote this shift in metabolism towards storage of nutrients (largely carbohydrates) as fat.

14 Our results are in apparent contrast with a previous publication showing that neonatal
15 ablation of Agrp neurons leads to decreased RER, increased metabolic efficiency and obesity in
16 adult animals ([Joly-Amado et al., 2012](#)). Contrary to adult ablation of Agrp neurons which leads
17 to cessation of feeding and death ([Gropp et al., 2005](#); [Luquet et al., 2005](#); [Wu and Palmiter,](#)
18 [2011](#)), neonatal ablation of these neurons is compatible with life ([Luquet et al., 2005](#)). In the
19 study by ([Joly-Amado et al., 2012](#)), Agrp neurons were ablated during the first postnatal week.
20 The authors found Agrp neuron-ablated mice presented metabolic abnormalities only later in
21 adulthood (> 3 months of age), which were in similar direction to acute activation of Agrp
22 neurons in adult animals (this report and ([Srisai, 2016](#); [Steculorum et al., 2016](#))). These results
23 strengthen the idea that Agrp neurons have distinct roles at different developmental stages

1 ([Dietrich et al., 2012](#)), and that altering the function of Agrp neurons early in life leads to
2 compensatory mechanisms with long lasting physiological implications.

3 Recent studies have described the *in vivo* dynamics of Agrp neuron activity ([Betley et al.,](#)
4 [2015](#); [Chen et al., 2015](#); [Mandelblat-Cerf et al., 2015](#)). However, not all cells were homogeneous
5 in their response. While 2/3 of Agrp neurons had decreased activity upon presentation of
6 food/eating, the other third did not change firing rate or even increased activity during eating.
7 The fact that Agrp neurons had heterogeneous response to feeding further highlights the
8 existence of subpopulations of Agrp neurons that are functionally, anatomically ([Betley et al.,](#)
9 [2013](#); [Padilla et al., 2016](#); [Steculorum et al., 2016](#)), and potentially genetically distinct. In light of
10 our data, it is thus possible that the sustained activity of distinct subpopulation of Agrp neurons
11 during feeding/refeeding might operate to maximize energy storage by shifting substrate
12 utilization towards lipogenesis.

13 Less expected than the elevated activity of Agrp neurons during food deprivation were
14 the findings that during diet-induced obesity (DIO) the activity of these neurons is also elevated
15 ([Baver et al., 2014](#); [Diano et al., 2011](#); [Dietrich et al., 2013](#); [Wei et al., 2015](#)). In a previous
16 study, we have identified that the elevated activity of Agrp neurons during high-fat feeding relies
17 on intracellular mitochondria fusion machinery ([Dietrich et al., 2013](#)). We found that deletion of
18 mitofusins selectively in Agrp neurons was sufficient to blunt the increase in neuronal activity in
19 response to a high-fat diet (HFD); as a consequence, mice were resistant to DIO. Intriguingly, we
20 found only minor or no effects of these genetic manipulations on food intake ([Dietrich et al.,](#)
21 [2013](#)), indicating that, in obesogenic conditions, Agrp neurons also control other aspects of
22 physiology that are independent of food consumption. The elevated activity of Agrp neurons
23 during obesity development ([Baver et al., 2014](#); [Diano et al., 2011](#); [Dietrich et al., 2013](#); [Wei et](#)

1 [al., 2015](#)) could be involved in the metabolic shifts towards fat deposition (lipogenesis) as
2 observed here. This consideration, however, will require further testing in future studies.

3 In summary, we showed that Agrp neurons in the hypothalamus rapidly control whole-
4 body nutrient utilization by shifting metabolism towards lipogenesis via the sympathetic nervous
5 system. These studies expanded the repertoire of functions attributed to Agrp neurons, which
6 emerge as controllers of a variety of physiological functions in addition to food intake. Because
7 there are several thousands of Agrp neurons in the mammalian brain, future studies will be
8 important to determine whether each subpopulation of Agrp neurons regulate a specific
9 physiological function. Because Agrp neurons release several molecules, including AGRP, NPY,
10 and GABA, it is also possible that this multifaceted function of Agrp neurons arises from the
11 combination of released molecules. These findings have strong implications for our
12 understanding of how neuronal circuits involved in metabolism regulation function and,
13 consequently, to our understanding of severe disordered conditions such as obesity.

1 **Experimental Procedures**

2 *Animals*

3 Mice used in the experiments were 3–8 months old from both genders. *Agrp*^{Trpv1} mice
4 were: *Agrp*^{Cre/+}::*Trpv1*^{KO/KO}::*R26-LSL-Trpv1*^{Gt/+}; control animals were either *Agrp*^{Trpv1} mice
5 injected with vehicle (3.3% Tween 80 in saline) or *Trpv1*^{KO/KO}::*R26-LSL-Trpv1*^{Gt/+} mice injected
6 with capsaicin. All animals were littermates (*Agrp* neuron activated and controls) in the
7 experiments. We did not observe any differences between the two control groups and, therefore,
8 throughout the manuscript we referred to them as “controls”. We have carefully characterized
9 this animal model to activate *Agrp* neurons and reported elsewhere ([Dietrich et al., 2015](#); [Ruan et](#)
10 [al., 2014](#)). We have performed dose-response curves for capsaicin and identified the dose of 10
11 mg/kg (i.p.) as optimal to induce behavior phenotypes in *Agrp*^{Trpv1} mice. We have also
12 performed a dose-response of capsaicin and measured changes in RER (1, 3, 10 and 30 mg/kg,
13 i.p.; experiments not reported). We also found that 10 mg/kg was the optimal dose to promote
14 changes in RER. Thus, we selected this dose of capsaicin for our studies.

15 The following mouse lines were used in this study: *Agrptm1*(cre)*Lowl/J*,
16 *Gt*(ROSA)26Sortm1(*Trpv1*,*ECFP*)*Mde/J*, *Trpv1tm1Jul/J*. These lines are available from The
17 Jackson laboratories. All animals were kept in temperature and humidity controlled rooms, in a
18 12/12h light/dark cycle, with lights on from 7:00AM-7:00PM. Food and water were provided ad
19 libitum, unless otherwise stated. All procedures were approved by IACUC (Yale University).

20 *Drugs*

21 The following compounds were used in the reported studies: C75 (RPMI medium 1640;
22 from Tocris); capsaicin (3.3% Tween-80 in saline; from Sigma), and CL-316, 245 (in PBS; from

1 Tocris). All drugs were injected in a volume of 10 ml/kg of body weight intraperitoneally (i.p.).
2 When multiple injections were performed in the same experiment, the volume of each injection
3 was adjusted to a total volume of 10 ml/kg per animal.

4 ***Metabolic Assays and Biochemical Analysis***

5 For all experiments animals were housed in individual cages at least three days prior to
6 the experiment. Blood samples were collected from the tail in order to measure glucose and free
7 fatty acids (NEFA) levels. Glucose was measured using a One Touch Ultra 2 glucometer. After
8 blood centrifugation, serum was collected and used to measure NEFA as indicated by
9 manufacturer (WAKO, Japan).

10

11 ***Gene Expression and Western blotting***

12 Animals were deeply anesthetized with ketamine and xylazine and euthanized by
13 decapitation. Tissues were collected and frozen in liquid nitrogen. Tissues were lysed in buffer
14 containing 1% Nonidet P-40, 50 mM Tris 3 HCl, 0.1 mM EDTA, 150 mM NaCl, proteinase
15 inhibitors and protein phosphatase inhibitors. Equal amounts of protein lysate were
16 electrophoresed on SDS-PAGE gels and transferred to PVDF membranes. Primary antibodies
17 (Lipolysis Activation Antibody Sampler Kit #8334, Cell Signaling) were incubated at 4°C
18 overnight. Membranes were washed and incubated with secondary antibodies conjugated to
19 horseradish peroxidase. Protein levels were visualized using ECL chemiluminescent substrate
20 and quantified using ImageJ.

21 Total RNA was extracted from mouse tissues using RNeasy® lipid mini kit (Qiagen).
22 cDNA was reverse transcribed (Bio-Rad) and amplified with SYBR Green Supermix (Bio-Rad)

1 using a Light Cycler 480 real-time PCR system (Roche). Data were normalized to the expression
2 of *Actin*, *Gusb* and *Arbp*. Primer sequences are available on request.

3 ***Indirect calorimetry***

4 Oxygen consumption (VO_2) and CO_2 production (VCO_2) were measured in four to eight
5 mice simultaneously in indirect calorimetry chambers (TSE Systems, Germany). Measurements
6 were recorded every 8-12 minutes over the entire course of the experiment (except for the
7 experiment in which ad libitum food intake was measured during one entire day). Respiratory
8 exchange ratio (RER) was calculated as the ratio between VCO_2 and VO_2 . Whole body fat
9 utilization was calculated using the follow equation: $1.67 * (VO_2 - VCO_2)$. Whole body
10 carbohydrate utilization was calculate using the follow equation: $4.55 * VCO_2 - 3.21 * VO_2$
11 (Frayn, 1983). All animals were single housed during the experiments in calorimetry chambers.
12 For the experiments in which different diets were fed to the animals, the following diets were
13 used: low-fat high-sugar diet (LFHS; D12450B, Open Source Diets, USA); high-fat diet 45%
14 calories from fat (HF45; D12451, Open Source Diets, USA); and high-fat diet 60% calories from
15 fat (HF60; D12492, Open Source Diets, USA). For the glucose response study, mice were
16 provided with glucose (D-Glucose, G8270, Sigma, USA) via gavage. We used saline as vehicle
17 and three doses of glucose (1, 2 and 3 g/kg body weight, via gavage feeding). Food was removed
18 2 hours before the experiment during the light cycle of the animals. Calorimetry was recorded
19 during 60 minutes prior injection of capsaicin and diet switch. For the experiments that no food
20 was provided, food was removed from the cages 2 hours before injecting mice with
21 capsaicin/vehicle. Baseline calorimetry was recorded for 60 minutes, and then the effects of
22 *Agrp* neuron activation were recorded for 60-120 minutes. Similar procedures were used in the
23 experiments in which compounds were given to mice prior the experiment. Glucose (2 g/kg, via

1 gavage), C75 (10 mg/kg, i.p.) or CL-316,243 (0.01-1.00 mg/kg, i.p) were given together with
2 capsaicin. In all cases, the drugs were injected immediately before capsaicin using two different
3 syringes. Total injected volume was adjusted for the maximal dose of 10 ml/kg mouse body
4 weight.

5 ***Statistical Considerations***

6 Matlab R2016a, PASW Statistics 18.0, Prism 7.0 and Adobe Illustrator CS6/CC were
7 used to analyze data and plot figures. Student's *t* test was used to compare two groups. ANOVA
8 was used to compare multiple groups. When necessary, multiple comparisons post hoc test
9 (MCT) was used (Holm-Sidak's test). When homogeneity was not assumed, the Kruskal–Wallis
10 nonparametric ANOVA was selected for multiple statistical comparisons. The Mann–Whitney U
11 test was used to determine significance between groups. Statistical data are provided in the
12 figures. $P < 0.05$ was considered statistically significant.

1 **Acknowledgments**

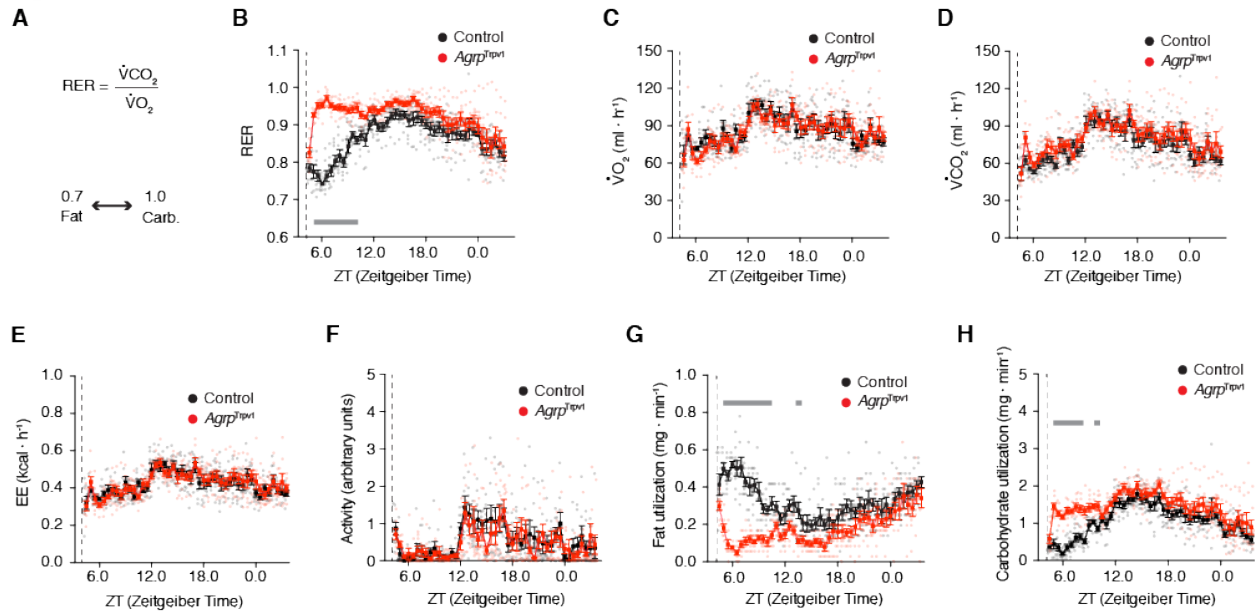
2 We thank Matthew Rodeheffer, Ivan de Araujo, Hai-bin Ruan and Xiaoyong Yang for comments
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10

11 **Author Contributions**

12 JB and MZ helped to perform the experiments and analyze the data. JAP and MOD performed
13 the experiments, designed, analyzed data and wrote the manuscript.

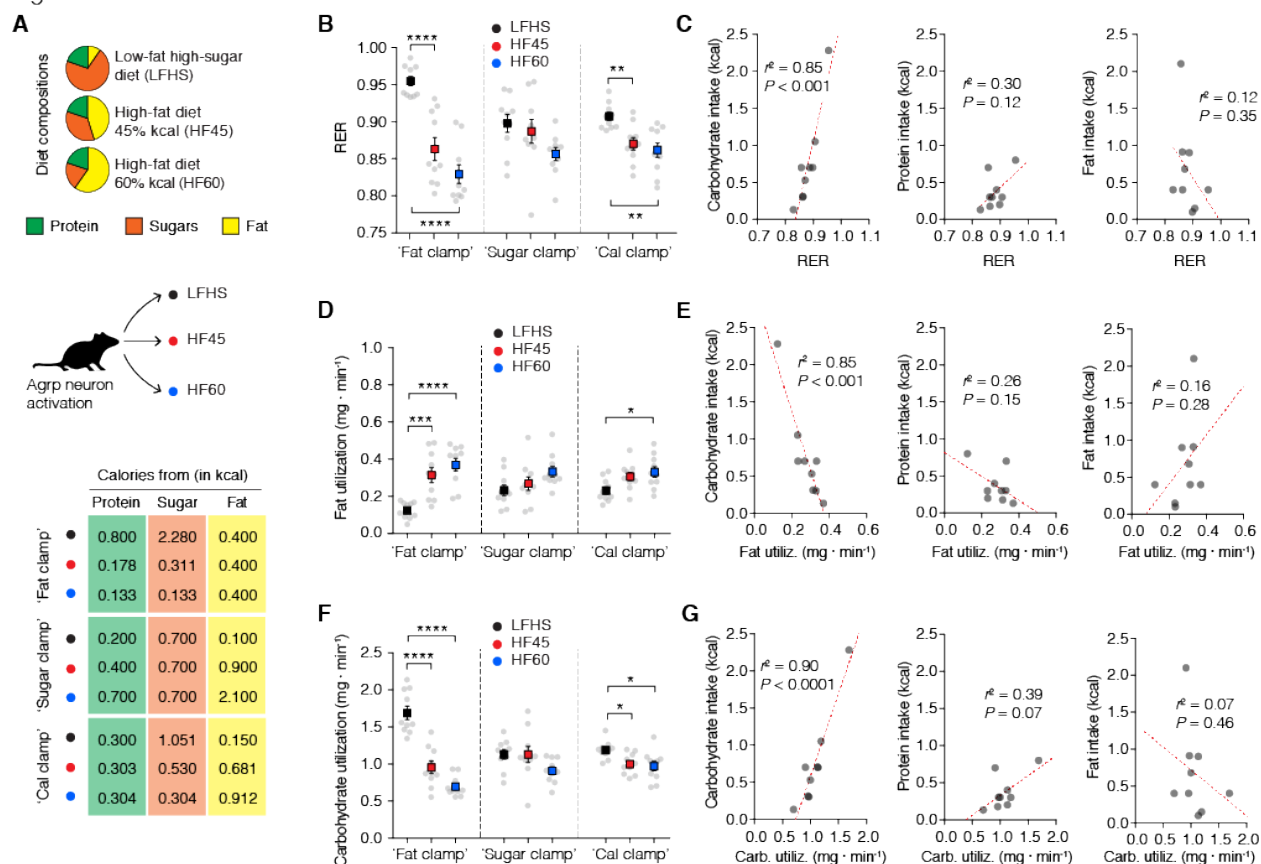
Figure 1



1
2 **Figure 1: Rapid shift in substrate utilization upon activation of AgRP neurons.**
3 (A) In indirect calorimetry chambers, RER was calculated by dividing the $\dot{V}CO_2$ by the $\dot{V}O_2$;
4 oxidation of fat acids (e.g., palmitate) generates a RER of 0.7, while oxidation of carbohydrates
5 (e.g., glucose) generates a RER of 1.0. From B-H, control (black; n = 8) and $AgRP^{Trpv1}$ mice (red;
6 n = 8) were acclimated to calorimetry chambers before been injected with capsaicin (dashed
7 lines; 10 mg/kg, i.p.) during the light cycle with food and water provided *ad libitum*. (B) RER
8 (interaction: $F_{45, 630} = 8.40, P < 0.0001$; time: $F_{45, 630} = 13.86, P < 0.0001$; group: $F_{1, 14} = 52.59, P$
9 < 0.0001). (C) $\dot{V}O_2$ (interaction: $F_{45, 630} = 1.07, P = 0.34$; time: $F_{45, 630} = 11.22, P < 0.0001$;
10 group: $F_{1, 14} = 0.01, P = 0.89$). (D) $\dot{V}CO_2$ (interaction: $F_{45, 630} = 0.88, P = 0.68$; time: $F_{45, 630} =$
11 $15.07, P < 0.0001$; group: $F_{1, 14} = 1.58, P = 0.22$). (E) Energy expenditure (interaction: $F_{45, 630} =$
12 $0.96, P = 0.53$; time: $F_{45, 630} = 12.09, P < 0.0001$; group: $F_{1, 14} = 0.02, P = 0.87$). (F) Ambulatory
13 activity (interaction: $F_{45, 630} = 0.80, P = 0.81$; time: $F_{45, 630} = 6.02, P < 0.0001$; group: $F_{1, 14} =$
14 $0.31, P = 0.58$). (G) Calculated fat utilization (interaction: $F_{45, 630} = 6.44, P < 0.001$; time: $F_{45, 630} =$
15 $7.22, P < 0.0001$; group: $F_{1, 14} = 33.83, P < 0.0001$). (H) Calculated carbohydrate utilization

1 (interaction: $F_{45, 630} = 2.77, P < 0.0001$; time: $F_{45, 630} = 18.48, P < 0.0001$; group: $F_{1, 14} = 16.73, P$
2 = 0.001). Statistical analysis was performed using two-way ANOVA with time as a repeated
3 measure followed by Holm-Sidak's multiple comparisons test (MCT). Grey bars indicate time
4 points in which MCTs were statistically significant ($P < 0.05$). Dashed line indicates time of
5 capsaicin injection. Small symbols indicate individual values. Large symbols indicate mean \pm
6 SEM.

Figure 2



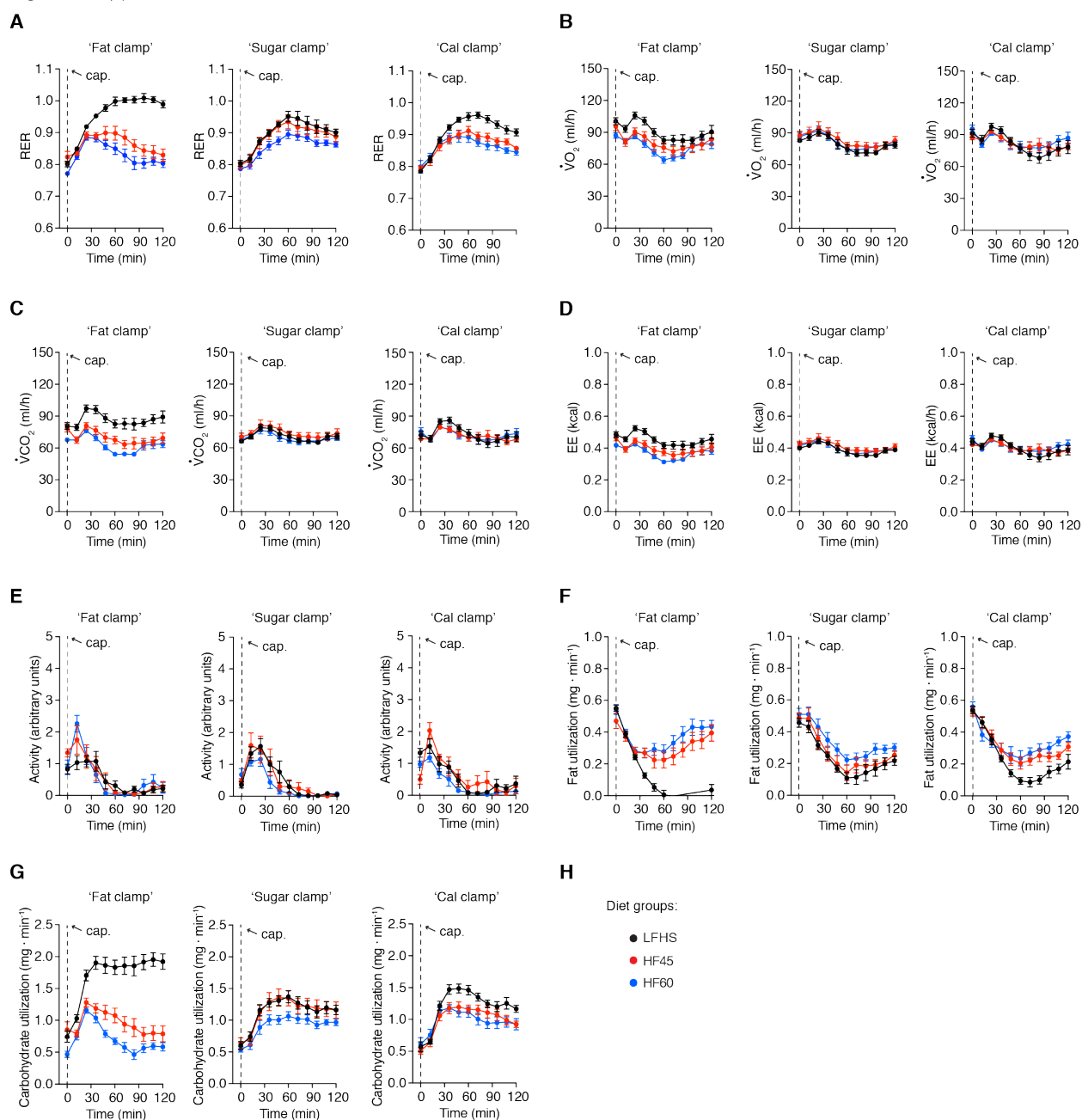
1

2 **Figure 2: Substrate utilization in response to different diets.**

3 Mice (*Agrp*^{Trpv1}) were tested in calorimetry chambers for their rapid response to different diets
4 upon activation of *Agrp* neurons (with capsaicin, 10 mg/kg, i.p). (A) Three different diets were
5 used in the studies, containing different distribution of macronutrients from similar sources
6 (protein levels were equal at 20 kcal%); low-fat high-sugar diet (LFHS; Research Diets
7 D12450B [carbohydrates: 70 kcal%; fat: 10 kcal%]); high-fat diet (HF45; Research Diets
8 D12451 [carbohydrates: 35 kcal%; fat: 45 kcal%]); and high-fat diet (HF60; Research Diets
9 D12492 [carbohydrates: 20 kcal%; fat: 60 kcal%]). Mice were fed pre-weighted amounts of the
10 three different diets in three different experimental conditions (see table; n = 10 mice per
11 condition): 'Fat clamp', all mice received a pellet of the diet containing 0.4 kcal from fat; 'Sugar
12 clamp', mice received a food pellet containing 0.7 kcal from carbohydrates; and 'Cal clamp',

1 mice received a food pellet containing a total of 1.5 kcal from all sources. **(B)** Mean RER after
2 activation of *Agrp* neurons (related to Figure 2 supplement 1A): ‘Fat clamp’ ($F_{2,27} = 29.13$, $P <$
3 0.0001); ‘Sugar clamp’ ($F_{2,27} = 2.662$, $P = 0.08$), and ‘Cal clamp’ ($F_{2,27} = 4.753$, $P = 0.01$). **(C)**
4 Correlation between RER (abscissa) and macronutrient intake (ordinate). **(D)** Mean calculated fat
5 utilization after activation of *Agrp* neurons (related to Figure 2 supplement 1F): ‘Fat clamp’ ($F_{2,27}$,
6 $27 = 16.5$, $P < 0.0001$); ‘Sugar clamp’ ($F_{2,27} = 2.878$, $P = 0.07$), and ‘Cal clamp’ ($F_{2,27} = 8.897$, P
7 $= 0.001$). **(E)** Correlation between fat utilization (abscissa) and macronutrient intake (ordinate).
8 **(F)** Mean calculated carbohydrate utilization after activation of *Agrp* neurons (related to Figure 2
9 supplement 1G): ‘Fat clamp’ ($F_{2,27} = 45.83$, $P < 0.0001$); ‘Sugar clamp’ ($F_{2,27} = 2.627$, $P =$
10 0.09), and ‘Cal clamp’ ($F_{2,27} = 5.53$, $P = 0.009$). **(G)** Correlation between carbohydrate
11 utilization (abscissa) and macronutrient intake (ordinate). In B, D and F, ordinary one-way
12 ANOVA was used to test for statistical differences within each experimental condition (‘Clamp’)
13 followed by Holm-Sidak’s multiple comparisons test (MCT). MCTs are indicated as * $P < 0.05$,
14 ** $P < 0.01$, *** $P < 0.001$, and **** $P < 0.0001$. Large symbols indicate mean \pm SEM. Small
15 grey symbols indicate individual values. Ten mice were used per condition. In C, E, and G,
16 linear regression analysis was used to calculate the correlation between total macronutrient
17 intake and substrate utilization measurements (as plotted in B, D, and F); each point represents
18 the mean response of 10 animals in the same condition (from B, D, and F); r^2 and P values are
19 plotted in each panel; dashed red line represents the linear regression model. Figure 2
20 supplement 1 presents detailed analysis of the calorimetry data and activity levels.

Figure 2 supplement 1



1

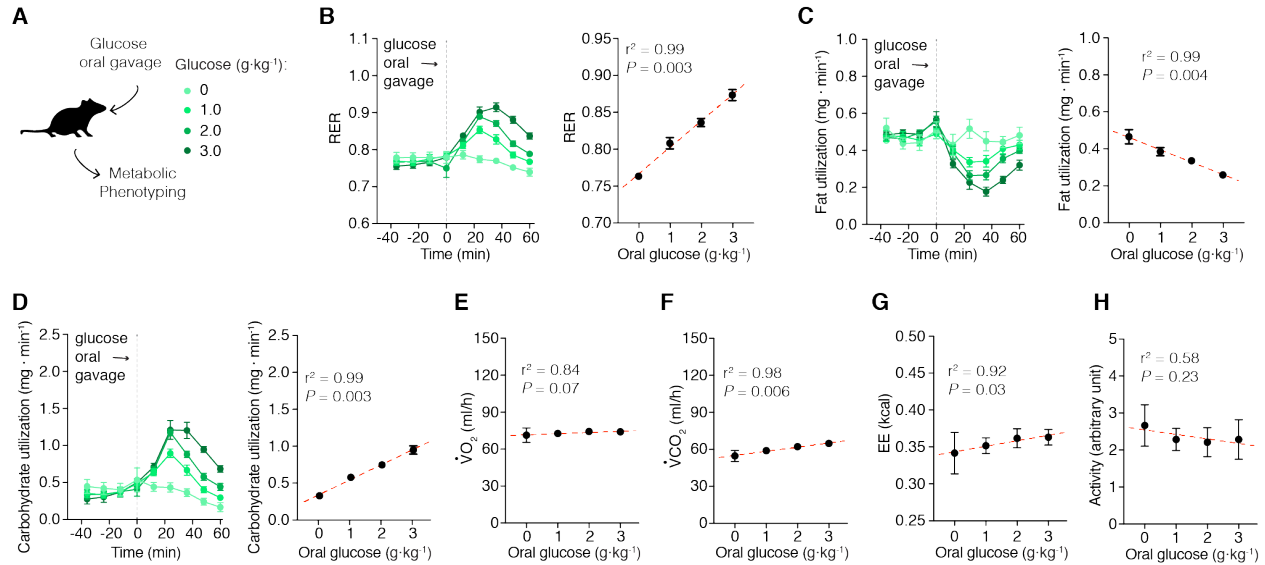
2 **Figure 2 supplement 1: Substrate utilization in response to different diets.**

3 Detailed metabolic phenotyping data of *Agrp*^{Trpv1} mice fed different diets (Figure 2A) upon
 4 activation of *Agrp* neurons by peripheral injection of capsaicin (10 mg/kg, i.p.). In black, mice
 5 fed a low-fat high-sugar diet (LFHS; Research Diets D12450B); in red, mice fed a high-fat diet
 6 (45 kcal% from fat, HF45; Research Diets D12451); and in blue, mice fed a high-fat diet (60

1 kcal% from fat, HF60; Research Diets D12492). (A) RER: ‘Fat clamp’ (interaction: $F_{20, 270} = 25$,
2 $P < 0.0001$; time: $F_{10, 270} = 56.34$, $P < 0.0001$; diets: $F_{2, 27} = 29.13$, $P < 0.0001$); ‘Sugar clamp’
3 (interaction: $F_{20, 270} = 0.64$, $P = 0.87$; time: $F_{10, 270} = 65.72$, $P < 0.0001$; diets: $F_{2, 27} = 2.87$, $P =$
4 0.07); ‘Cal clamp’ (interaction: $F_{20, 270} = 3.51$, $P < 0.0001$; time: $F_{10, 270} = 49.89$, $P < 0.0001$;
5 diets: $F_{2, 27} = 8.92$, $P = 0.001$). (B) VO_2 : ‘Fat clamp’ (interaction: $F_{20, 270} = 0.91$, $P = 0.56$; time:
6 $F_{10, 270} = 16.08$, $P < 0.0001$; diets: $F_{2, 27} = 6.28$, $P < 0.0001$); ‘Sugar clamp’ (interaction: $F_{20, 270} =$
7 0.18 , $P > 0.99$; time: $F_{10, 270} = 14.66$, $P < 0.0001$; diets: $F_{2, 27} = 0.69$, $P = 0.50$); ‘Cal clamp’
8 (interaction: $F_{20, 270} = 1.65$, $P = 0.04$; time: $F_{10, 270} = 13.98$, $P < 0.0001$; diets: $F_{2, 27} = 0.13$, $P =$
9 0.87). (C) VCO_2 : ‘Fat clamp’ (interaction: $F_{20, 270} = 1.70$, $P = 0.03$; time: $F_{10, 270} = 11.55$, $P <$
10 0.0001 ; diets: $F_{2, 27} = 22.47$, $P < 0.0001$); ‘Sugar clamp’ (interaction: $F_{20, 270} = 0.28$, $P = 0.99$;
11 time: $F_{10, 270} = 8.59$, $P < 0.0001$; diets: $F_{2, 27} = 0.90$, $P = 0.41$); ‘Cal clamp’ (interaction: $F_{20, 270} =$
12 1.30 , $P = 0.17$; time: $F_{10, 270} = 10.33$, $P < 0.0001$; diets: $F_{2, 27} = 0.14$, $P = 0.86$). (D) Energy
13 expenditure: ‘Fat clamp’ (interaction: $F_{20, 270} = 0.84$, $P = 0.66$; time: $F_{10, 270} = 14.62$, $P < 0.0001$;
14 diets: $F_{2, 27} = 8.92$, $P = 0.001$); ‘Sugar clamp’ (interaction: $F_{20, 270} = 0.19$, $P > 0.99$; time: $F_{10, 270}$
15 $= 12.9$, $P < 0.0001$; diets: $F_{2, 27} = 0.67$, $P = 0.51$); ‘Cal clamp’ (interaction: $F_{20, 270} = 1.55$, $P =$
16 0.06 ; time: $F_{10, 270} = 12.66$, $P < 0.0001$; diets: $F_{2, 27} = 0.08$, $P = 0.91$). (E) Ambulatory activity:
17 ‘Fat clamp’ (interaction: $F_{20, 270} = 1.69$, $P = 0.03$; time: $F_{10, 270} = 25.32$, $P < 0.0001$; diets: $F_{2, 27} =$
18 0.23 , $P = 0.78$); ‘Sugar clamp’ (interaction: $F_{20, 270} = 1.17$, $P = 0.27$; time: $F_{10, 270} = 31.81$, $P <$
19 0.0001 ; diets: $F_{2, 27} = 1.01$, $P = 0.37$); ‘Cal clamp’ (interaction: $F_{20, 270} = 1.68$, $P = 0.03$; time: $F_{10,$
20 $270} = 26.55$, $P < 0.0001$; diets: $F_{2, 27} = 3.03$, $P = 0.06$). (F) Fat utilization: ‘Fat clamp’
21 (interaction: $F_{20, 270} = 19.24$, $P < 0.0001$; time: $F_{10, 270} = 63.42$, $P < 0.0001$; diets: $F_{2, 27} = 16.5$, P
22 < 0.0001); ‘Sugar clamp’ (interaction: $F_{20, 270} = 0.34$, $P = 0.99$; time: $F_{10, 270} = 55.74$, $P < 0.0001$;
23 diets: $F_{2, 27} = 2.65$, $P = 0.08$); ‘Cal clamp’ (interaction: $F_{20, 270} = 3.22$, $P < 0.0001$; time: $F_{10, 270} =$

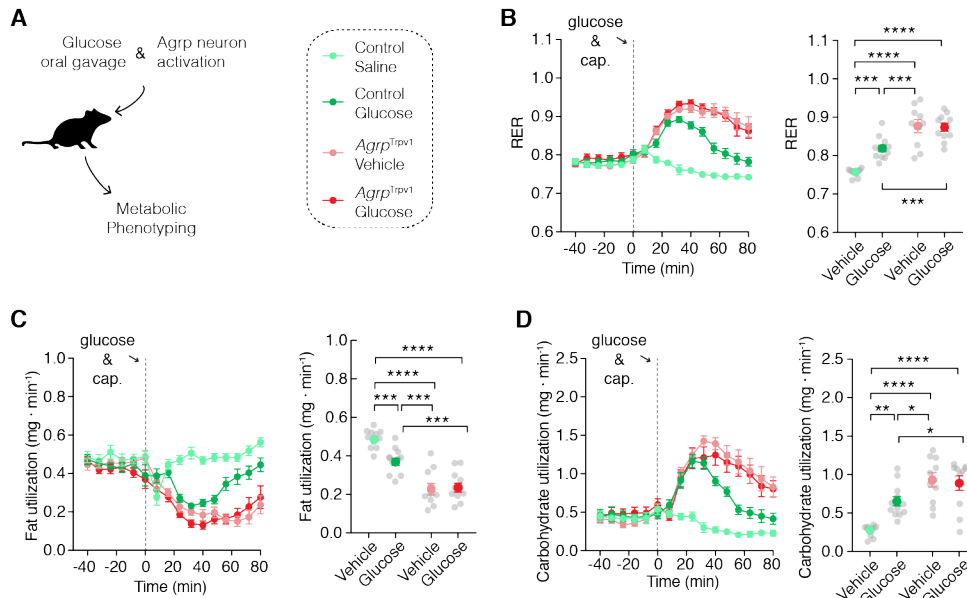
1 55.86, $P < 0.0001$; diets: $F_{2,27} = 4.76$, $P = 0.01$). (G) Carbohydrate utilization: ‘Fat clamp’
2 (interaction: $F_{20,270} = 14.97$, $P < 0.0001$; time: $F_{10,270} = 30.32$, $P < 0.0001$; diets: $F_{2,27} = 45.82$, P
3 < 0.0001); ‘Sugar clamp’ (interaction: $F_{20,270} = 0.67$, $P = 0.84$; time: $F_{10,270} = 44.44$, $P < 0.0001$;
4 diets: $F_{2,27} = 2.62$, $P = 0.09$); ‘Cal clamp’ (interaction: $F_{20,270} = 2.13$, $P = 0.003$; time: $F_{10,270} =$
5 44.43 , $P < 0.0001$; diets: $F_{2,27} = 5.52$, $P = 0.009$). Statistical analysis was performed using two-
6 way ANOVA with time as a repeated measure followed by Holm-Sidak’s multiple comparisons
7 test (MCT). MCTs are not shown. Dashed line indicates time of capsaicin injection. A total n =
8 10 mice were used for each condition. Symbols indicate mean \pm SEM.

Figure 3



1
2 **Figure 3: Acute effects of glucose on metabolism.**
3 Metabolic phenotyping of mice immediately after a dose-response of glucose (via gavage). (A)
4 Mice received a bolus of saline or glucose solution (1, 2, or 3 g/kg body weight dissolved in
5 saline) via gavage. (B) Glucose gavage infusion produces an acute increase in RER that is dose
6 dependent. (C) Calculated fat utilization negatively correlates to glucose ingestion. (D)
7 Calculated carbohydrate utilization is positively correlated to glucose ingestion. (E) $\dot{V}O_2$ levels
8 did not correlated with glucose infusion. (F) $\dot{V}CO_2$ was positively correlated with glucose
9 ingestion. (G) Glucose gavage also positively correlated with energy expenditure measurements,
10 but did not correlated with changes in (H) activity levels. The total number of animals used in
11 this study was: saline (n = 7); glucose 1 g/kg (n = 12); glucose 2 g/kg (n = 16); and glucose 3
12 g/kg (n = 8). Symbols represent mean \pm SEM. Grey dashed line indicates time of oral gavage. In
13 the linear correlation panels, symbols indicate mean of all mice in the given group \pm SEM;
14 dashed red line represents the linear regression model; r^2 and P values are plotted in each panel.

Figure 4



1

2 **Figure 4: Agrp neurons control substrate utilization independently of ingestion.**

3 (A) Control and *Agrp*^{Trpv1} mice received a bolus of saline or glucose (2 g/kg) via gavage

4 followed by peripheral injection of capsaicin (10 mg/kg, i.p.). (B) Changes in RER in control

5 mice fed different doses of glucose solutions (interaction: $F_{1,40} = 9.82$, $P = 0.003$; gavage

6 solution: $F_{1,40} = 8.20$, $P = 0.006$; genotype: $F_{1,40} = 71.31$, $P < 0.0001$). (C) Fat utilization

7 (interaction: $F_{1,40} = 7.91$, $P = 0.007$; gavage solution: $F_{1,40} = 6.07$, $P = 0.01$; genotype: $F_{1,40} =$

8 79.82 , $P < 0.0001$). (D) Carbohydrate utilization: (interaction: $F_{1,40} = 8.29$, $P = 0.006$; gavage

9 solution: $F_{1,40} = 5.26$, $P = 0.02$; genotype: $F_{1,40} = 37.44$, $P < 0.0001$). Statistical analysis was

10 performed using two-way ANOVA on the mean response after gavage and capsaicin injection;

11 genotype (control vs. *Agrp*^{Trpv1}) and gavage infusion (saline vs. glucose) were used as factors for

12 the ANOVA. Holm-Sidak's multiple comparisons test (MCT) was used to find post-hoc

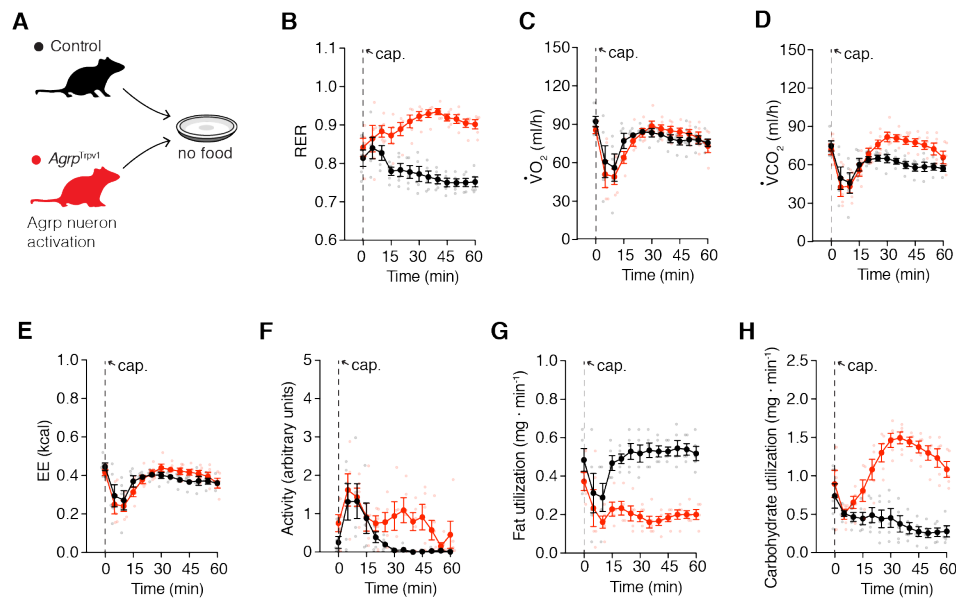
13 differences among groups. MCTs are indicated as * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$, and

14 **** $P < 0.0001$ in figure panels. Control mice + saline gavage (n = 11); control mice + glucose

15 gavage (n = 12); *Agrp*^{Trpv1} mice + saline gavage (n = 10); *Agrp*^{Trpv1} mice + glucose gavage (n =

- 1 11). Dashed grey line indicates time of oral gavage and capsaicin injection. Colored symbols
- 2 indicate mean \pm SEM. Grey symbols indicate individual values.

Figure 4 supplement 1



1

2 **Figure 4 supplement 1: Agrp neurons control substrate utilization independently of**

3 **ingestion.**

4 (A) Well-fed control (in black) and *Agrp*^{Trpv1} (in red) mice were tested in metabolic chambers

5 upon injection of capsaicin without food provided. (B) RER (interaction: $F_{12,96} = 8.96$, $P <$

6 0.0001 ; time: $F_{12,96} = 1.31$, $P = 0.22$; genotype: $F_{1,8} = 54.45$, $P < 0.0001$). (C) $\dot{V}O_2$ (interaction:

7 $F_{12,96} = 0.87$, $P = 0.57$; time: $F_{12,96} = 9.63$, $P < 0.0001$; genotype: $F_{1,8} = 0.15$, $P = 0.70$).

8 (D) $\dot{V}CO_2$ (interaction: $F_{12,96} = 2.89$, $P = 0.001$; time: $F_{12,96} = 10.66$, $P < 0.0001$; genotype: $F_{1,8}$

9 $= 10.39$, $P = 0.01$). (E) Energy expenditure (interaction: $F_{12,96} = 1.16$, $P = 0.31$; time: $F_{12,96} =$

10 9.89 , $P < 0.0001$; genotype: $F_{1,8} = 0.13$, $P = 0.72$). (F) Ambulatory activity (interaction: $F_{12,96} =$

11 1.12 , $P = 0.34$; time: $F_{12,96} = 7.13$, $P < 0.0001$; genotype: $F_{1,8} = 5.29$, $P = 0.05$). (G) Calculated

12 fat utilization (interaction: $F_{12,96} = 3.52$, $P = 0.0002$; time: $F_{12,96} = 3.23$, $P = 0.0006$; genotype:

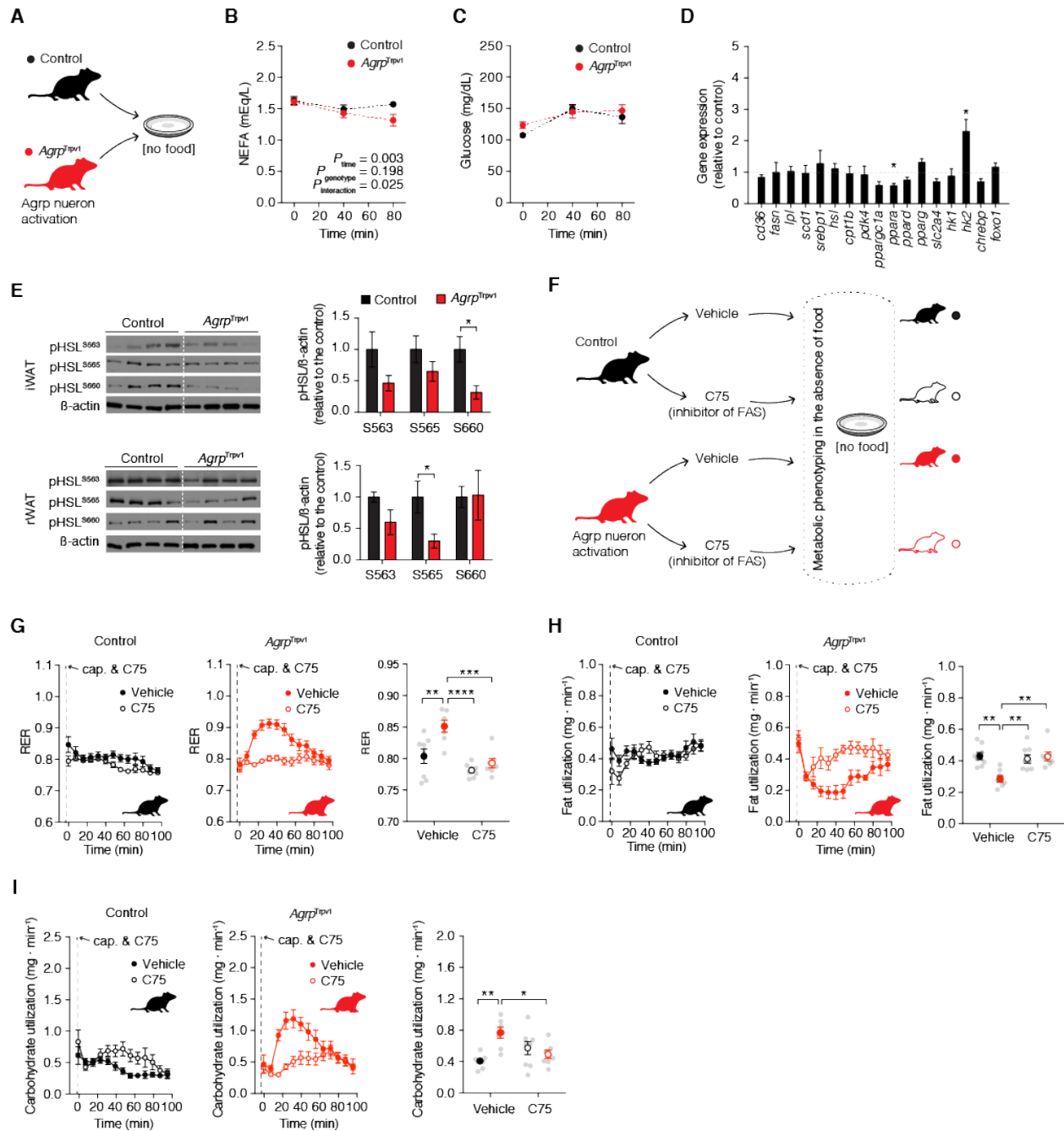
13 $F_{1,8} = 55.31$, $P < 0.0001$). (H) Calculated carbohydrate utilization (interaction: $F_{12,96} = 13.52$, P

14 < 0.0001 ; time: $F_{12,96} = 6.25$, $P < 0.0001$; genotype: $F_{1,8} = 58.16$, $P < 0.0001$). Statistical

15 analysis was performed using two-way ANOVA with time as a repeated measure followed by

- 1 Holm-Sidak's multiple comparisons test (MCT). MCTs are not shown. Dashed line indicates
- 2 time of capsaicin injection. Small symbols indicate individual values. Large symbols indicate
- 3 mean \pm SEM.

Figure 5



1

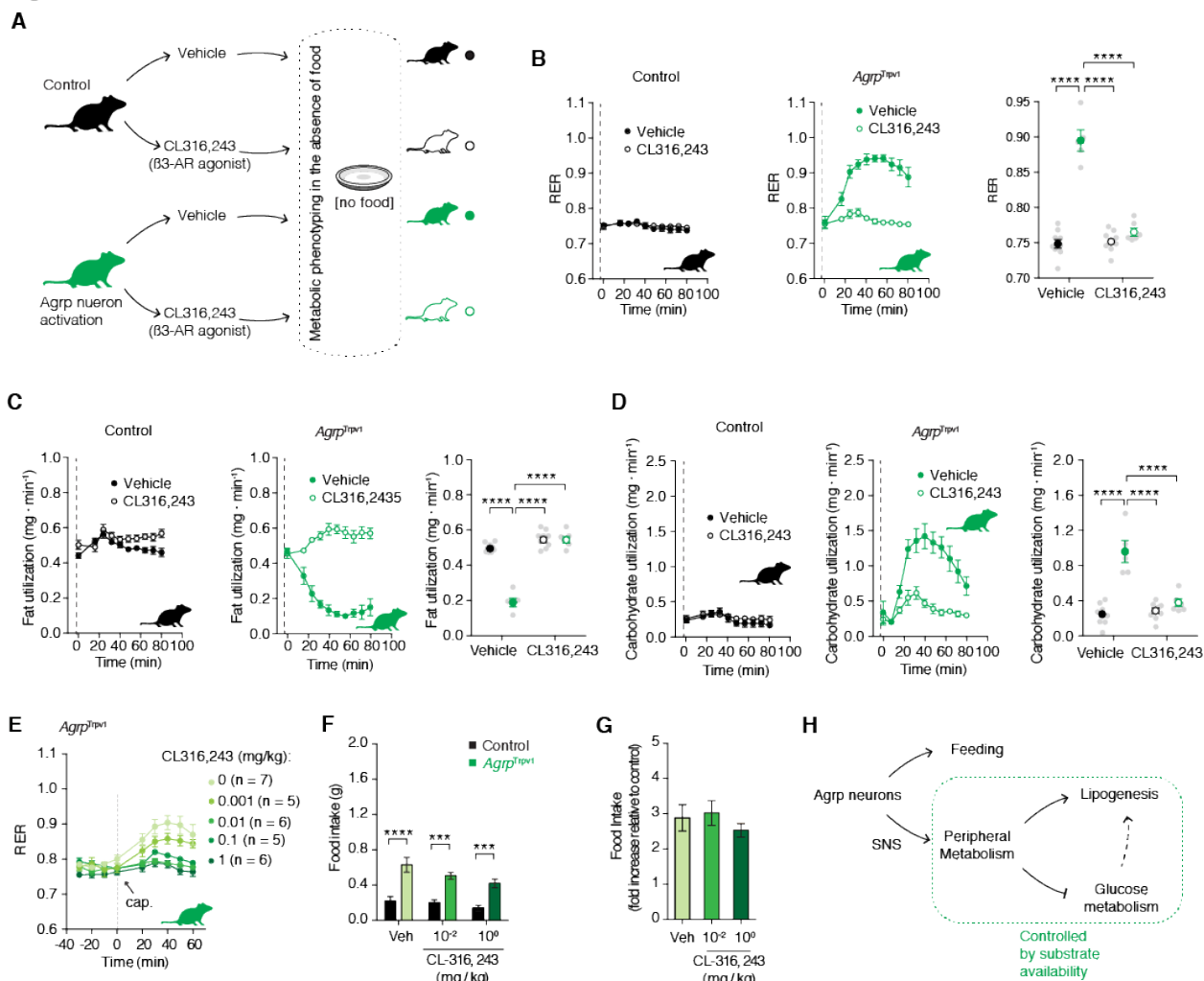
2 **Figure 5: Agrp neurons promote lipogenesis.**

3 (A) Well-fed control (in black) and *Agrp^{Trpv1}* (in red) mice were injected with capsaicin (10
4 mg/kg, i.p.). One group of animals was tail bled before and after (40 and 80 min) injection to
5 measure blood levels of NEFA and glucose. Different groups of animals were sacrificed 60

1 minutes after capsaicin injection and white adipose tissue (WAT) was collected for biochemical
2 analyses. **(B)** NEFA levels (interaction: $F_{2, 54} = 3.91$, $P = 0.02$; time: $F_{2, 54} = 9.64$, $P = 0.0003$;
3 genotype: $F_{1, 27} = 1.74$, $P = 0.19$; MCT: $P_{80 \text{ min}} = 0.03$). **(C)** Blood glucose levels (interaction: $F_{2,$
4 $54 = 1.47$, $P = 0.23$; time: $F_{2, 54} = 14.6$, $P < 0.0001$; genotype: $F_{1, 27} = 0.67$, $P = 0.41$; MCT: not
5 significant). In B and C: control (n = 15) and *Agrp*^{Trpv1} mice (n = 14). Statistical analysis was
6 performed using two-way ANOVA with time as a repeated measure followed by Holm-Sidak's
7 multiple comparisons test (MCT). **(D)** Analysis of the transcriptional profile of the WAT upon
8 activation of *Agrp* neurons. Data are normalized to control levels (dashed line). Statistical
9 analysis was performed using student's *t*-test. * $P < 0.05$. **(E)** Western blotting analysis of
10 phosphorylated HSL in inguinal WAT and retroperitoneal WAT (n = 4 mice per group).
11 Statistical analysis was performed using student's *t*-test. * $P < 0.05$. **(F)** Experimental design to
12 test the participation of fatty acid synthetase (FAS) in the effects of *Agrp* neurons on substrate
13 utilization. Control and *Agrp*^{Trpv1} mice were randomized to receive vehicle or the FAS inhibitor
14 (C75, 10 mg/kg, i.p.) immediately before capsaicin injection. **(G)** RER (interaction: $F_{1, 26} = 4.36$,
15 $P = 0.04$; drug: $F_{1, 26} = 22.28$, $P < 0.0001$; genotype: $F_{1, 26} = 11.73$, $P = 0.002$). **(H)** Fat
16 utilization (interaction: $F_{1, 26} = 9.97$, $P = 0.04$; drug: $F_{1, 26} = 6.00$, $P = 0.02$; genotype: $F_{1, 26} =$
17 6.56 , $P = 0.01$). **(I)** Carbohydrate utilization: (interaction: $F_{1, 26} = 12.18$, $P = 0.001$; drug: $F_{1, 26} =$
18 0.76 , $P = 0.38$; genotype: $F_{1, 26} = 4.90$, $P = 0.03$). In G-I, statistical analysis was performed using
19 two-way ANOVA on the mean response after drug and capsaicin injection; genotype (control vs.
20 *Agrp*^{Trpv1}) and drug (vehicle vs. C75) were used as factors for the ANOVA. Holm-Sidak's
21 multiple comparisons test (MCT) was used to find post-hoc differences among groups. MCTs are
22 indicated as * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$, and **** $P < 0.0001$ in figure panels.
23 Control mice + vehicle (n = 8); control mice + C75 (n = 8); *Agrp*^{Trpv1} mice + vehicle (n = 7);

- 1 *Agrp*^{Trpv1} mice + C75 (n = 7). Dashed line indicates time of injections. Colored symbols indicate
- 2 mean ± SEM. Grey symbols indicate individual values.

Figure 6



1

2 **Figure 6: SNS signaling is involved in peripheral effects of AgRP neurons.**

3 (A) Experimental design to test the participation of the sympathetic nervous system (SNS) in the

4 effects of AgRP neurons on substrate utilization. Control (black) and *AgRP^{Tripv1}* mice (green) were

5 randomized to receive vehicle or the β3-adrenergic receptor agonist (CL316,243, 1 mg/kg, i.p.);

6 vehicle or CL316,243 were injected immediately before capsaicin. (B) RER (interaction: $F_{1,25} =$

7 75.09, $P < 0.0001$; drug: $F_{1,25} = 67.96$, $P < 0.0001$; genotype: $F_{1,25} = 108.4$, $P < 0.0001$).

8 (C) Fat utilization (interaction: $F_{1,25} = 85.56$, $P < 0.0001$; drug: $F_{1,25} = 150.4$, $P < 0.0001$; genotype: $F_{1,$

9 25 = 86.42, $P < 0.0001$). (D) Carbohydrate utilization: (interaction: $F_{1,25} = 29.74$, $P < 0.0001$;

10 drug: $F_{1,25} = 22.75$, $P < 0.0001$; genotype: $F_{1,25} = 50.12$, $P < 0.0001$). In B-D, statistical analysis

1 was performed using two-way ANOVA on the mean response after drug and capsaicin injection;
2 genotype (control vs. *Agrp*^{Trpv1}) and drug (vehicle vs. CL316,243) were used as factors for the
3 ANOVA. Holm-Sidak's multiple comparisons test (MCT) was used to find post-hoc differences
4 among groups. MCTs are indicated as * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$, and **** $P <$
5 0.0001 in figure panels. Control mice + vehicle (n = 9); control mice + CL316,243 (n = 9);
6 *Agrp*^{Trpv1} mice + vehicle (n = 5); *Agrp*^{Trpv1} mice + CL316,243 (n = 6). Dashed line indicates time
7 of injections. Colored symbols indicate mean \pm SEM. Grey symbols indicate individual values.
8 (E) Dose-response of CL316,243 injected immediately before capsaicin in *Agrp*^{Trpv1} mice
9 (dashed line denotes injection time). Number of animals used per experimental group shown in
10 the panel. (F) Food intake response of control and *Agrp*^{Trpv1} mice to activation of *Agrp* neurons
11 when injected with different doses of CL316,243 (n = 7 for all groups); interaction ($F_{2, 36} = 0.95$,
12 $P = 0.39$); drug ($F_{2, 36} = 4.16$, $P = 0.02$); genotype ($F_{1, 36} = 64.96$, $P < 0.0001$). Statistical analysis
13 was performed using two-way ANOVA with genotype and drug as factors. Holm-Sidak's
14 multiple comparisons test (MCT) was used to find post-hoc differences genotypes. MCTs are
15 indicated as *** $P < 0.001$ and **** $P < 0.0001$ in figure panel. (G) Related to F, but fold
16 change in food intake in *Agrp*^{Trpv1} related to control mice. Bars and symbols indicate mean \pm
17 SEM. (H) Diagram illustrating the model for different roles of *Agrp* neurons in the control of
18 feeding and peripheral metabolism. *Agrp* neurons control peripheral metabolism via the SNS, but
19 switching substrate utilization towards use of carbohydrate.

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