Ciliary gene **RPGRIP1L** is required for hypothalamic arcuate neuron development

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**Intronic polymorphisms in the α-ketoglutarate–dependent dioxygenase gene (**FTO**)** that are highly associated with increased body weight have been implicated in the transcriptional control of a nearby ciliary gene, retinitis pigmentosa GTPase regulator-interacting protein-1 like (**RPGRIP1L**). Previous studies have shown that congenital **Rpgrip1l** hypomorphism in murine proopiomelanocortin (Pomc) neurons causes obesity by increasing food intake. Here, we show by congenital and adult-onset **Rpgrip1l** deletion in Pomc-expressing neurons that the hyperphagia and obesity are likely due to neurodevelopmental effects that are characterized by a reduction in the Pomc/Neuropeptide Y (Npy) neuronal number ratio and marked increases in arcuate hypothalamic–paraventricular hypothalamic (ARH-PVH) axonal projections. Biallelic **RPGRIP1L** mutations result in fewer cilia-positive human induced pluripotent stem cell–derived (iPSC-derived) neurons and blunted responses to Sonic Hedgehog (SHH). Isogenic human ARH-like embryonic stem cell–derived (ESC-derived) neurons homozygous for the obesity-risk alleles at **rs8050136** or **rs1421085** have decreased **RPGRIP1L** expression and have lower numbers of POMC neurons. **RPGRIP1L** overexpression increases POMC cell number. These findings suggest that apparently functional intronic polymorphisms affect hypothalamic **RPGRIP1L** expression and impact development of POMC neurons and their derivatives, leading to hyperphagia and increased adiposity.

**Introduction**

Genome-wide association studies have identified common obesity variants that map at over 100 loci across the genome (1). Our aim is to deconvolute the strongest genetic signal for adiposity within an ~47-kb interval in the first intron of α-ketoglutarate–dependent dioxygenase gene (**FTO**). Specific noncoding alleles within this interval are associated with ~3.5-pound and ~2-pound body weight differences, per risk allele, in adults and children, respectively (2–6). Two of the associated single nucleotide polymorphisms...
(SNPs), rs8050136 and rs1421085, alter DNA-binding of CUX1 (7, 8), a transcription factor involved in neurodevelopment and highly expressed in the arcuate hypothalamus (ARH) (7, 9, 10). In the CNS, CUX1 regulates the expression of FTO and 5′-adjacent RPGRIP1L (7, 8, 11) encoding a protein located at the transition zone of the primary cilium (12). Gene mutations in components of the primary cilium cause syndromic and nonsyndromic forms of obesity, such as Bardet-Biedl Syndrome (BBS), Alström, tubby (13–15), and MC4R/AC3 (16). The subcellular localization of RPGRIP1L at the gate between cytoplasm and cilium suggests it may regulate entry and exit of ciliary signaling components relating to energy balance. In support of this hypothesis, the leptin receptor long isoform (Lepr-b) colocalizes with RPGRIP1L (17). We have also reported that Rpgrip1l expression is downregulated in the hypothalamus of leptin-deficient mice and is upregulated by leptin administration (11), indicating that RPGRIP1L may convey anorectic effects via the canonical leptin signaling pathway. Mice with congenital systemic deletion of 1 Rpgrip1l copy, or 2 Rpgrip1l copies in leptin-receptor isoform b-expressing neurons, are hyperphagic, obese, and display diminished suppression of food intake in response to leptin administration (8, 17). Hence, it is possible that RPGRIP1L modulates food intake in adults through cilia-dependent regulation of Lepr-b trafficking and function.

Embryonic lethality due to Rpgrip1l global deletion in mice is accompanied by severe CNS abnormalities of the cerebellar vermis and midbrain, as well as dilation of ventricles (18, 19). Consistent with phenotypic effects in the CNS, Rpgrip1l hypomorphism in the developing cerebral cortex impairs neuronal migration (20). In humans, RPGRIP1L mutations in Joubert Syndrome (JBST) cause “the molar tooth” characterized by cerebral vermian hypoplasia, accompanied by elongation of the caudal tegmentum and dysplasia of the caudal medulla (19). The severity of developmental defects in individuals who lack functional RPGRIP1L may mask a potential role in energy homeostasis. However, a recent report described a JBST patient with mutations in ciliary adenosine diphosphate (ADP) ribosylation factor–like GTPase 13B (ARL13B) who presented with obesity and structural brain abnormalities similar to those individuals with RPGRIP1L inactivating mutations (21), suggesting that anatomical/functional CNS abnormalities caused by RPGRIP1L haploinsufficiency may cause obesity in humans.

Loss of RPGRIP1L in human primary cell cultures and mice results in increased ciliary axonemal length (17, 22), and systemic overexpression of Cux1 in mice increases cilia length and causes polycystic kidney disease, a typical ciliopathic phenotype (23). Collectively, these reports suggest that RPGRIP1L and CUX1 may be part of the same cilia-related pathway. Moreover, CUX1 loss of function affects aspects of cortical neurocircuit assembly of the corpus callosum (9, 10). Given the observed CNS and ciliary anatomical changes caused by alterations in RPGRIP1L and CUX1 expression, we hypothesized that CUX1-dependent transcriptional control of RPGRIP1L may account for some of the association of FTO intronic SNPs with adiposity by modulating regulation of food intake via cilia-dependent alterations of hypothalamic structure.

**Results**

**Induction of Rpgrip1l hypomorphism in the adult ARH causes obesity**

*Arcuate stereotaxic injections.* Mice segregating for 1 congenital null Rpgrip1l allele systemically (Rpgrip1l<sup>−/−</sup>) ingest more regular chow as early as 4 weeks of age and display increased adiposity by 12 weeks of age (17). Just as in congenital Rpgrip1l hypomorphism, Rpgrip1l hypomorphism induced in the adult CNS using tamoxifen-induced Nestin-Cre results in increased body weight and increased fat mass associated with increased food intake (8). To further refine the brain regions responsible for the increased food intake, we stereotaxically injected Cre recombinase-expressing adeno-associated viral particles (Cre-AAV) or GFP-expressing AAV (GFP-AAV) into the ARH of 12-week-old male Rpgrip1l<sup>−/−</sup> mice (Figure 1A). The mouse cohorts injected in the ARH were also compared with littermates that were targeted with Cre-AAV in both the dorsomedial (DMH) and ventromedial (VMH) hypothalami. By 16 weeks of age, ARH expression of Rpgrip1l in Cre-AAV–injected mice was decreased by ~90%; these mice were ~50% heavier than animals in which Cre-AAV had been injected in the DMH + VMH or mice GFP-AAV–injected in the ARH (Figure 1, B and C). The increase in body weight in mice injected with Cre-AAV in the ARH compared with mice injected with GFP-AAV in the ARH was due to an ~3-fold increase in fat mass (Figure 1, C and D). These findings suggest that effects of Rpgrip1l on body weight are conveyed, in part, via the ARH in adult animals and that Rpgrip1l hypomorphism induced in the DMH + VMH of adult animals has no impact on adiposity.
Cerebellar stereotaxic injections. Because RPGRIP1L mutations in JBST patients result in cerebellar vermis hypoplasia/aplasia (24), we stereotaxically injected Cre-AAV into the developing cerebellar vermis of P10 male Rpgrip1fl/fl mice (Figure 2A). At P10, the cerebellar vermis is still under development (25). Similar to the DMH + VMH, Rpgrip1l hypomorphism in the cerebellar vermis did not increase adiposity in mice fed regular chow or a high-fat diet (Figure 2B). These data indicate that cerebellar Rpgrip1l may not be implicated in the regulation of food intake.

Adult-onset Rpgrip1l hypomorphism specifically in Pomc neurons does not cause adiposity

We have reported that 16-week-old mice congenitally hypomorphic for Rpgrip1l in Pomc-expressing neurons (Rpgrip1l−/−[Pomc]) are obese due to increased food intake (8). However, tamoxifen-induced loss of Rpgrip1l in terminally differentiated Pomc neurons in adult animals (Rpgrip1l−/−[adult Pomc-CreER]) at 8 weeks of age did not cause significant increases in body weight (Figure 3). In contrast, systemic Rpgrip1l loss (Rpgrip1l−/−[adult Cagg-CreER]) induced in adults phenocopies the obesity observed in Rpgrip1l−/−[Pomc] mice (Figure 3). These findings suggest that the developmental impact of Rpgrip1l hypomorphism on Pomc neurons is the sole driver for the increased adiposity in Rpgrip1l−/−[Pomc] mice.

Congenital Rpgrip1l hypomorphism in Pomc neurons alters Pomc and Npy neuron number

Pomc-GFP–positive adult neuron number assessed by IHC. We hypothesized that congenital Rpgrip1l−/−[Pomc] mice may have reduced numbers of POMC ARH neurons. In 10 μm coronal sections spanning the ARH of adult male Rpgrip1l−/−[Pomc] mice, we found that the number of GFP-positive ARH neurons is reduced by ~50% (Figure 4A).

Cell death of Pomc neurons assessed by TUNEL. We examined the possibility that Rpgrip1l deletion in Pomc neurons leads to increased cell death. By TUNEL staining, we were unable to detect increased cell death in the developing hypothalamus of Rpgrip1l−/−[Pomc] E11 embryos (when Pomc expression is initiated) or by induced Rpgrip1l deletion in Nestin-expressing adult hypothalamic neurons (Supplemental Figure 1; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.123337DS1). These findings suggest that congenital Rpgrip1l hypomorphism does not cause cell death.

Pomc progenitor neuron number assessed by iDISCO. Using the brain-clearing iDISCO system and IHC, we quantified the number of Pomc-expressing ARH neurons of Rpgrip1l−/−[Pomc] mice at E13 when the numbers of Pomc ARH progenitors are in highest (26, 27). We noted an ~20% decrease in Pomc-expressing ARH cells in Rpgrip1l−/−[Pomc] mice (Figure 4, B and C).

Pomc ARH-PVH projections assessed by IHC. We also utilized the mT/mG allele (28) in which the membrane-bound tdTomato (mT) marker is globally expressed in all tissues except those Pomc-Cre–expressing cells in which recombination results in silencing of mT and expression of membrane-bound GFP (mGFP), allowing visualization of cell bodies, projections, and terminals. We noted a marked increase in the density of anterior medial hypothalamic projections to the paraventricular hypothalamus (PVH) in Rpgrip1l−/−[Pomc] and Rpgrip1lfl/fl(Pomc) mice (Figure 4, D and E).

Npy-positive adult neuron number assessed by IHC. Bbs2- or Bbs6-null obese mice (murine models of the Bardet-Biedl ciliopathy) have an ~20% reduction in mature Pomc ARH neurons, as assessed by an anti-Pomc antibody (29). In contrast, in our congenital Rpgrip1l−/−[Pomc] mice, the Pomc-GFP transgene marks mature Pomc neurons that actively express GFP, as well as ARH neurons derived from Pomc-expressing early developmental progenitors (e.g., Npy neurons; refs. 26, 27). Pomc is expressed by the majority of cells in the developing ventral hypothalamus at E10–E11 and is silenced later in gestation in more than half of these cells, some of which become Npy neurons and some of which are directed to other terminal fates (26, 27). Therefore, the adiposity and hyperphagia of congenital Rpgrip1l−/−[Pomc] mice may be due to developmental changes in both mature Pomc ARH neurons and Pomc-derived ARH neurons. We explored the possibility that Rpgrip1l hypomorphism in Pomc neurons affected the number of Npy-expressing neurons that derive from Pomc-expressing progenitors. The total number of Npy-GFP–positive ARH neurons is increased by ~12% in Rpgrip1l−/−[Pomc] adult mice due to a 2-fold increase in the number of Npy neurons in the rostral ARH of Rpgrip1l−/−[Pomc] mice (Figure 5). Respective increases and decreases of hypothalamic Pomc and Npy neurons have been implicated in hypophagia (30, 31). The converse changes in the congenital Rpgrip1l−/−[Pomc] mice are consistent with their hyperphagia.
Diminished ciliation of iPSC-derived neurons of patients with RPGRIP1L mutations

Generation of iPSCs from JBST patients. We and others have demonstrated that Sonic Hedgehog (SHH) signaling activation promotes human stem cell differentiation to ARH-type neurons that predominantly express POMC (70%-80%) and exhibit other hypothalamus-specific patterns of gene transcription (32–34). We generated iPSCs from three JBST patient primary skin fibroblast lines UW15 (35), UW04-03 (36), and UW04-04 (17) with biallelic mutations in the RPGRIP1L C2 domain that severely disrupt — or are predicted to disrupt — nephrocystin-4 binding that facilitates recruitment of other proteins to the base of the cilium (37). The iPSCs expressed the expected pluripotency markers (Supplemental Figure 2). Pluripotency was further confirmed by injection of the iPSCs into immunocompromised NOD scid \(\gamma\) (NSG) mice resulting in the formation of teratomas that display ectodermal, mesodermal, and endodermal structures (Supplemental Figure 3A). The derived iPSCs had normal karyotypes (Supplemental Figure 3B).

Generation of JBST neurons from iPSCs. We attempted to generate ARH-like neurons from the JBST iPSCs but were unable to generate NKX2.1-positive neuronal progenitors from which POMC and NPY neurons derive (38, 39), further suggesting the importance of RPGRIP1L in the development of ARH neurons. Subsequently, we utilized a dual-SMAD inhibition protocol (40) to direct differentiation of the JBST iPSCs into neurons positive for neuromarkers Nestin, neuron-specific class III \(\beta\)-tubulin (TUJ1), and Neurofilament (NF) (Supplemental Figure 3C).

Figure 1. Adult-onset ARH Rpgrip1l deletion increases adiposity. (A) Stereotaxic injections of Cre-GFP–expressing AAV in the ARH (Rpgrip1l\(^{fl/fl}\);Cre-GFP;ARH) or Cre-expressing AAV particles in the DMH and VMH (Rpgrip1l\(^{fl/fl}\);Cre-GFP;DMH+VMH) of male mice. Scale bars: 1 mm. (B) Decreased Rpgrip1l expression in the ARH of Rpgrip1l\(^{fl/fl}\);Cre-GFP;ARH (n = 6) and in the DMH/VMH of Rpgrip1l\(^{fl/fl}\);Cre-GFP;DMH+VMH mice (n = 5). Rpgrip1l expression is indistinguishable in the ARH, DMH, or VMH of mice stereotaxically injected with GFP-expressing in the ARH (Rpgrip1l\(^{fl/fl}\);GFP;ARH; n = 6). P < 0.05 was statistically significant, by 1-way ANOVA. (C) Increased body weight and fat mass in Rpgrip1l\(^{fl/fl}\);Cre-GFP;ARH mice. P < 0.05 was statistically significant, by 1-way ANOVA. (D) Mass increase of various fat depots of Rpgrip1l\(^{fl/fl}\);Cre-GFP;ARH mice compared with Rpgrip1l\(^{fl/fl}\);GFP;ARH mice. Data in B, C, and D are represented as box-and-whisker plots; boxes are the interquartile range, lines are the median value, and whiskers are minimum and maximum values. *P < 0.01, by 2-tailed Student’s t test.
Ciliation in iPSC-derived JBST neurons. ARL13B is a small GTPase that participates in ciliary formation and function (41). About 60% of control neurons and ~40% of JBST neurons were ARL13B-positive (Figure 6, A–C). ARL13B, IFT20, and IFT88 mRNA levels were indistinguishable between control and JBST neurons (Figure 6D), suggesting that the ~30% decrease in ARL13B-positive JBST neurons is due to lack of cilia. No differences were observed in ciliary length between JBST and control neurons (Figure 6E).

RPGRIP1L hypomorphism is associated with perturbed SHH signaling

SHH conveys important hypothalamic neurospecification signals in vitro (32); these are mediated via the primary cilium (39, 42). When SHH binds to its receptor, PATCHED1, the liganded receptor exits the primary cilium with GFP-expressing AAV particles (Rpgrip1fl/fl;Cre-GFP;VERMIS; n = 8). Scale bar: 100 μm. (B) No difference in total body weight, fat mass, or lean mass between Rpgrip1fl/fl;Cre-GFP;VERMIS and Rpgrip1fl/fl;GFP;VERMIS male mice on ad lib regular chow up to 7 weeks of age or after 2 weeks on high-fat diet. Data in A are represented as box-and-whisker plots; boxes are the interquartile range, lines are the median value, and whiskers are minimum and maximum values. *P < 0.01, 2-tailed Student’s t test.

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SHH signaling in iPSC-derived JBST neurons. In JBST human fibroblasts, SMO failed to localize to the primary cilium upon SHH exposure (Figure 7B). In iPSC-derived JBST neurons, AC3 expression was decreased by ~60% and was ectopically localized in the neuron cell body rather than the
primary cilia upon SHH stimulation (Figure 7C). Finally, upregulation of PATCHED1 and GLI1 expression by ~40% and ~90%, respectively, after SHH treatment in control iPSC-derived neurons, was blunted in iPSC-derived JBST neurons (Figure 7D), further suggesting aberrant SHH signaling.

Alleles at rs1421085/rs8050136 and RPGRIPL expression correlate with human embryonic stem cell–derived (ESc-derived) POMC-positive ARH neuron number

Obesity-associated SNPs in FTO intron 1, rs8050136 and rs1421085, are embedded within 2 binding sites for CUX1 (8). Based on electrophoretic mobility shift assays (7, 11), the rs8050136 A (risk) and rs1421085 C (risk) alleles preferentially bind CUX1 isoform P200, whereas rs8050136 C (protective) and rs1421085 T (protective) alleles preferentially bind CUX1 isoform P110. Luciferase-based promoter probing of the rs8050136 and rs1421085 genomic regions in hypothalamus-derived N2a cells, and systemic overexpression of Cux1 isoforms p200 and p110 in vivo (8, 17), implicate P200 as a repressor of Fto and P110 as an activator of RPGRIPL expression in the CNS.

Alteration of rs8050136 and rs1421085 by CRISPR. Using the human H9 ESc line that is heterozygous for rs8050136 and rs1421085 risk alleles, we have used Clustered Regularly Interspaced Short Palindromic Repeats (CRISPR)/CRISPR-associated protein-9 (Cas9) technology to create an isogenic allelic series for the adiposity risk (R) or protective-associated (P-associated) alleles at SNP rs8050136 or rs1421085 in human ESc–derived ARH neurons (Supplemental Figure 4A) (45). We have confirmed that these lines are correctly targeted, karyotypically normal, and display no apparent off-target effects (45).

FTO/RPGRIPL expression per rs8050136/rs1421085 allele in isogenic ARH neurons. Homozygosity for the risk allele at rs8050136 or rs1421085 resulted in respective ~45% and ~75% decreases in generation of ARH-type neurons compared with neurons homozygous for the protective allele at rs8050136 or rs1421085 (Figure 8A). Homozygosity for the risk allele at rs8050136 or rs1421085 correlated with an ~20%–30% decrease in FTO or RPGRIPL expression in ARH-type neurons (Figure 8B).

POMC neuron number per rs8050136/rs1421085 allele in isogenic ARH neurons. Homozygosity for the risk allele at rs8050136 or rs1421085 was associated with an ~30% decrease in POMC-positive neurons (Figure 8C) compared with neurons homozygous for the protective allele at rs8050136 or rs1421085.
FTO/RPGIP1L expression per rs8050136/rs1421085 allele in isogenic ARH neurons overexpressing P200. We overexpressed CUX1 P200 at day 26 of the differentiation protocol, which is 1 day before differentiating cells normally initiate expression of POMC (32, 34) (Claudia Doege, Columbia University, personal communication). P200 overexpression by lentiviral transduction of ARH neurons homozygous for the risk allele (preferentially bound by P200) at rs8050136 or rs1421085 decreased FTO expression by ~60% (Figure 8B). P200 overexpression also resulted in increased P110 protein levels (Supplemental Figure 4, B and C), presumably due to constitutive Cathepsin L processing of P200–P110 (7). Increased P110 protein levels in neurons homozygous for the risk alleles at rs8050136 or rs1421085 were associated with an ~20%–30% increase in RPGRIP1L expression (P = 0.06), presumably due to the lower DNA-binding affinity of P110 for the risk alleles (7).

POMC neuron number per rs8050136/rs1421085 obesity-risk allele in isogenic ARH neurons overexpressing P200. The increase in RPGRIP1L expression in ARH neurons homozygous for the risk alleles at rs8050136 or rs1421085 overexpressing P200 was associated with restoration of POMC-positive neuron number (Figure 8C). Furthermore, RPGRIP1L overexpression (~2-fold) by lentiviral transduction of neurons homozygous for the obesity-risk allele at rs1421085 was associated with an ~50% increase in the number of POMC-positive neurons (Figure 8C).
Fto does not affect Pomc neuron number. Because (a) RPGRIP1L overexpression alone is sufficient to increase POMC neuron number and (b) the increase in RPGRIP1L expression and decreased FTO expression upon P200 overexpression restores POMC number in neurons homozygous for the risk alleles at rs8050136 or rs1421085, it is implied that RPGRIP1L is important for POMC neuron specification and that FTO protein does not participate in the differentiation of POMC neurons. In support of the latter inference, mice homozygous for the Fto–floxed allele (Ftofl/fl) (8) and also segregating for Pomc-Cre and Pomc-GFP have no apparent change in the number of adult GFP-positive neurons compared with Ftofl/fl littermates (Supplemental Figure 5).

**Discussion**

The hyperphagia and obesity caused by induced Rpgrip1l deletion in the terminally differentiated CNS (8) (Figure 1) supports the importance of Rpgrip1l in the maintenance and/or neurochemical function of the developed feeding neurocircuitry. It is likely that diminished leptin sensitivity is partly responsible for the increased energy intake. The fact that adult-onset Rpgrip1l deletion in Pomc neurons did not result in increased body weight implies that Rpgrip1l is not essential for the neurochemical function of Pomc adult neurons. Instead, the data presented here support a critical role of RPGRIP1L in the development of Pomc-expressing neuroprogenitors, including Npy neurons that derive from Pomc-expressing progenitors (26, 27). The reciprocal decrease in anorexigenic Pomc neurons and increase in orexigenic Npy neurons may provide a cellular anlage for the hyperphagia of Rpgrip1l–/–(Pomc) mice. Our finding that POMC neuron specification can be altered by modulating RPGRIP1L expression per genotype at obesity risk alleles at rs1421085 or rs8050136 supports the relevance of this mechanistic formulation to the adiposity highly associated with allelic variation at these FTO intronic loci.
The Pomm-GFP transgene that we used also necessarily marks those neurons that develop from Pomm-expressing neuroprogenitors. We anticipate that specific neuronal subpopulations are affected by Pomm-specific congenital deletion of Rpgrip1l. However, with the exception of Npy neurons — some of which are known to develop from Pomm precursors — we have not identified these neuronal subpopulations. Therefore, the precise neurocellular consequences of the congenital interference with Rpgrip1l expression in Rpgrip1l–/–(Pomc) mice are not known. Future work should be directed toward identifying all the ARH neuronal subpopulations affected in Rpgrip1l–/–(Pomc) mice, as well as assigning the affected ARH-PVH projections to specific ARH neuronal subpopulations.

The increase in PVH projections emanating from the ARH of Rpgrip1l–/–(Pomc) mice may partly reflect the increase in the number of Npy neurons. However, the large magnitude of the increase in projections, and the relatively smaller magnitude in increase of Npy number in Rpgrip1l–/–(Pomc) mice, suggests that other ARH neurons that derive from Pomm-expressing progenitors are also affected. Alternatively, Rpgrip1l hypomorphism may relieve inhibition of axon growth/guidance (46) leading to this apparent change in wiring of the feeding neurocircuitry.

Using a hypothalamic differentiation protocol reported elsewhere (32, 34), the human H9 ESc line and “CRISPERed” isogenic lines produced 70%–80% POMC-positive neurons, whereas less than 1% of ARH neurons were NPY positive. By E18, more than half of POMC neurons convert to NPY-positive neurons during embryogenesis (26). We believe that the differentiation protocol employed here gives rise to early
POMC progenitors that may resemble those at E13 in the mouse in vivo. The decreased POMC cell number in the human ESc–derived ARH neurons segregating for the obesity risk alleles at rs1421085 or rs8050136 is in agreement with the reduction in Pomc neuroprogenitors in Rpgrip1l–/–(Pomc) embryos at stage E13.

We were unable to detect any increase in cell death in Rpgrip1l–/–(Pomc) or control embryos at E11 when Pomc expression turns on in the developing hypothalamus; cell death in the ARH is not common at this developmental stage (Lori Zeltser, Columbia University; personal communication). Similarly, adult-onset Rpgrip1l deletion in the CNS using Nestin-Cre did not increase cell death in the adult ARH. It is possible that we have missed a narrow time window at which increased apoptosis may have occurred in Rpgrip1l–/–(Pomc) neurons. Ex vivo live imaging of Pomc progenitors at the developmental stage when POMC expression is normally initiated might provide insights into this issue. However, the fact that Npy neuron number increased in the adult ARH of Rpgrip1l–/–(Pomc) mice suggests a cell specification–related phenotype. Moreover, the human fibroblasts with biallelic mutations in RPGIRP1L used in this study proliferate at normal rates, suggesting that perturbed RPGRIP1L function does not promote apoptosis.

An important question is whether RPGRIP1L hypomorphism causes proliferative and/or cell fate specification changes in POMC-expressing progenitors or neurons that derive from them. RPGRIP1L, a component

Figure 7. Sonic Hedgehog (SHH) signaling is impaired in JBST patient fibroblasts and neurons. (A) IHC showing decreased Smo-positive Pomc-expressing neuroprogenitors (green, Pomc-GFP) in E13 Rpgrip1l–/–(Pomc) (n = 3) compared with Rpgrip1lf/f embryos (n = 3). Scale bars: 10 μm. (B) Immunostaining of Smoothened (SMO) and Acetylated-tubulin (Ac-Tub) showing that JBST fibroblasts fail to translocate SMO to the cilia upon SHH stimulation (100 ng/ml). Scale bar: 5 μm. Arrow indicates primary cilium. (C) Diminished AC3 expression and localization in the ciliary axoneme of iPSC-derived JBST neurons in the absence of SHH. SHH stimulation resulted in ectopic AC3 localization in the neuron cell body in IPSC-derived JBST neurons. Neuron cell body and projections are also marked with neuro-specific markers Neurofilament (NF) and NeuN. Each column represents the average of 3 cell lines. Scale bar: 100 μm. (D) PATCHED1 and GLI1 expression is not increased in iPSC-derived neurons from JBST patients UW04-03 and UW04-04 upon SHH stimulation. Each column represents the average of 3 cell lines. Expression was normalized to the TATA box binding protein (TBP). *P < 0.01, by 2-tailed Student’s t test.
of the centrosome, and CUX1 are implicated in the control of the cell cycle (47, 48), a cellular process coordinated with ciliary formation arising from the centrosome and ciliary disassembly (49). Nevertheless, it is not clear whether the primary cilium regulates cell division. On the other hand, the importance of the primary cilium in the control of SHH signaling and neuropatterning is well established (42). Shh signaling is disrupted in systemic Rpgrip1l-null embryos in which CNS structural abnormalities are prominent (18, 50). We conclude that RPGRIP1L regulates Pomc neuron specification by a SHH-dependent mechanism.

The primary cilium is also involved in cell migration. The platelet-derived growth factor receptor α (PDGFrα) localizes to the primary cilium and is part of the PDGF signaling pathway that controls cell migration during development and in the adult via AKT signaling and mammalian target of rapamycin complex 1 (mTORC1) pathways (51, 52). RPGRIP1L has been implicated in the mTORC1 pathway (53), and Rpgrip1l hypomorphism results in perturbed neuronal migration of the murine developing cortex (20). The potential effects of Rpgrip1l hypomorphism on Pomc-neuron migration during development, as well as potential effects on neuronal function of Pomc neurons and other neuronal subpopulations affected, are areas of interest.

Pomc is expressed in other sites implicated in feeding behavior, such as the pituitary and brain stem (54, 55), and Pomc neurons in the ARH project to other brain regions, such as the dorso vagal complex (55).
Therefore, Rpgrip1l hypomorphism may affect Pmc number and projections in other regions in a manner similar to the ARH.

A recent study indicates that the melanocortin 4 receptor (MC4R) is located on the primary cilium and that hypomorphic Mc4r mutations impair the receptor’s ciliary trafficking in PVH neurons (56). In the same study, inhibition of ciliary Ac3 in Mc4r-expressing neurons was associated with obesity. Shh reduces cAMP levels in the primary cilium by inhibiting adenylyl cyclase activity (57). We observed decreased AC3 (ADCY3) expression and ectopic AC3 localization upon SHH stimulation in JBST iPSC–derived neurons. It is plausible that upregulation of AC3 expression in response to adenylyl cyclase inhibition by Shh in circumstances of Rpgrip1l hypomorphism results in accumulation of AC3 in the cell body due to the deranged AC3 trafficking. However, we find that Rpgrip1l hypomorphism suppresses adiposity in Ac3−/− mice, suggesting that Rpgrip1l is functionally downstream of AC3 and that the AC3-mediated adiposity is not caused by Rpgrip1l hypomorphism (8). Therefore, it is unlikely that decreased AC3 expression and ectopic AC3 localization are driving the developmental defects caused by Rpgrip1l hypomorphism in Pmc neurons. On the other hand, AC3 and cAMP signaling mediate axon outgrowth and guidance in the CNS (58) and may be, at least in part, responsible for the perturbed Pmc neuron axonal outgrowth in circumstances of Rpgrip1l deficiency.

Previous studies have produced conflicting results regarding the correlation of allelic variations at the FTO intronic region with FTO expression or expression of nearby genes. Smemo et al. (59) reported higher expression of iroquois homeobox 3 (IRX3) in human cerebellar and adipose tissue homozygous for FTO obesity risk alleles at rs9930506 compared with tissue from individuals homozygous for the protective FTO alleles. In contrast, partial inhibition of Irx3 in the murine hypothalamus has the opposite effect of what would be predicted by the direction of the association or by Irx3 global deletion (59, 60). Moreover, Claussnitzer et al. (61) failed to see an allele-specific effect on IRX3 expression in a hippocampal neuronal cell line, but they reported effects of allelic variants at rs1421085 on IRX3 and -5 (IRX3/5) in preadipocytes. Claussnitzer et al. proposed a mechanism by which obesity risk alleles of rs1421085 in human mesenchymal adipocyte progenitors increase expression of IRX3/5, resulting in decreased thermogenic potential of new adipocytes. However, a follow-up study presented in vivo and in vitro evidence that Irx3 positively regulates thermogenesis (62). The relevance of these mechanisms in humans is unclear, as the effects of the obesity-risk alleles in the first intron of FTO have been inconsistently reported to convey effects on food intake and not energy expenditure (6, 63–71). The inconsistencies in correlation of allelic variation to expression of FTO and vicinal genes may be attributed to the possibility that different haplotypes within intron 1 affect different genes and, thereby, different physiological process. This possibility would be consistent with multiple mechanisms (72) at different levels of penetrance and within different race/ethnicities, underlying such a statistically strong effect of noncoding alleles on adiposity. The present study focuses on the action of Rpgrip1l in neurons, in which expression of Rpgrip1l, but not IRX3/IRX5, is correlated in an FTO intronic allele dose-dependent manner (8).

In summary, we present evidence that the very strong association with adiposity of highly prevalent sequence variants in intron 1 of FTO are due, in part, to their indirect impact on the primary cilium. Our data imply that genetic mediation of nonsyndromic, prevalent obesity is conveyed partly by structural alteration of the hypothalamic cells and circuitry that mediate aspects of ingestive behaviors.

**Methods**

**Mouse studies.** Rpgrip1l+/−, Pmc-Cre (The Jackson Laboratory; stock number 005965), Pmc-GFP (The Jackson Laboratory; stock number 009593), and Nestin-CreER72 mice (The Jackson Laboratory; stock number 016261) have been described by us elsewhere (8). The Cagg-CreER allele (Tg[CAG-cre/Esr1]5Amc/J, The Jackson Laboratory, stock number 004682) used to assess systemic Rpgrip1l loss in adults was induced as previously reported (73). For adult-induced loss of Rpgrip1l specifically in adult terminally differentiated POMC neurons, a POMC-CreER allele that has previously been described (74) was utilized. mT/mG mice were purchased from The Jackson Laboratory (B6.129[Cg]-Gt[ROSA]26Sortm4[ACTB-tdTomato,−EGFP]Luo/J; stock number 007676). Npy-GFP mice were also purchased from The Jackson Laboratory (Npy-GFP [C57BL/6J-Tg[NPY-hrGFP]; stock number 006417]). Fto+− mice are described elsewhere (8). Mice where housed in a Barrier facility at room temperature (23°C) on a 12-hour light/12-hour dark cycle (lights off at 7 p.m.), with ad libitum (ad lib) access to food and water.
**Human studies.** UW15 (7), UW04-04 (17), and UW04-03 (36) primary skin fibroblast JBST lines were provided by Daniel Doherty (University of Washington, Seattle, Washington, USA). ESc line H9 was purchased by WiCell Research Institute Inc. (catalog WA09).

**Diet and dietary treatment.** All mice were fed regular chow (Purina Mills; Picolab, catalog 5058) (16) ad lib unless specified. High-fat diet consists of 65% of calories as fat (catalog D12492; Open Source Diets).

**Body mass and composition measurements.** Measurements were taken with a Minispec TD-NMR Analyzer (Bruker Optics) as described elsewhere (8).

**Isolation of total RNA and nuclear protein extracts.** Total RNA and protein extraction have been described elsewhere (17). Specific brain regions were micropunched as described in Stratigopoulos et al. (7).

**Quantitative PCR (qPCR) and cDNA synthesis.** The method has been described elsewhere (7). The geometric mean of GAPDH and ACTB expression levels was used as loading controls, unless specified. The PCR primers used are shown in Supplemental Table 1.

**IHC, immunofluorescence, and antibodies.** IHC was performed on cryosections after perfusion fixation, as described elsewhere (17). Teratomas were embedded in paraffin and further cut into 8 μm-thick sections. Sections were stained with H&E using a standard protocol (http://www.ihcworld.com/protocols/special_stains/h&e_ellis.htm). Cells growing on cultured-cell slides pretreated with Poly-L-ornithine/laminin were fixed with 4% paraformaldehyde for 10 minutes at room temperature. Primary fibroblasts were starved for 36 hours before visualization of primary cilia by immunofluorescence. The following primary antibodies were used: anti-GFP (1:100; Abcam, catalog ab6662), TUJ1 (1:500; MilliporeSigma, catalogs T2200 and T5076), MAP2 (1:10,000; Abcam, catalog AB5392), AC3 (1:100; Santa Cruz Biotechnology Inc., catalog sc-588), ARL13B (1:1,000; gift from Tamara Caspary, Emory University, Atlanta, Georgia, USA), SMO (1:500; gift from Kathryn V. Anderson, Memorial Sloan Kettering Cancer Center, New York, New York, USA), NF (1:10,000; MilliporeSigma, catalog AB5539), γ-tubulin (1:500; MilliporeSigma, catalog T6557), NESTIN (1:500; Stemgent, catalog 09-0045), NeuN (1:100; MilliporeSigma, catalog MAB377), Acetylated tubulin (1:500; MilliporeSigma, catalog T6793), POMC (1:500; Abcam, catalog ab14064), NPY (1:500; Novus, catalog nbp1-19808), αMSH (1:500; MilliporeSigma, catalog AB5087), and Gad67 (1:500; MilliporeSigma, catalog MAB5406).

**TUNEL.** TUNEL staining was performed using the DeadEnd Fluorometric TUNEL System (Promega, catalog G3250) according to the manufacturer’s instructions.

**Scanning electron microscopy.** iPSC-derived neurons were grown on 13 mm glass coverslips and fixed in 2.5% glutaraldehyde in 0.1 M Cacodylate buffer for 30 minutes at room temperature. Sample preparation and imaging for scanning electron microscope was done by the analytical imaging facility at Albert Einstein College of Medicine (New York, New York, USA). Experimental details are described elsewhere (75). Images were obtained at 5000× magnification.

**Brain clearing and imaging.** For brain clearing, we followed the protocol described in great detail at https://idisco.info/idisco-protocol/. Briefly, signal was enhanced by treatment with anti-GFP antibody (Aves Laboratories, catalog GFP-1020) and secondary Alexa Fluor 647 (anti-chicken; Thermo Fisher Scientific, catalog A-21449), followed by dehydration performed with tert-Butanol. Delipidization was achieved by treatment with dichloromethane (DCM). Then, samples were stored overnight in Benzyl alcohol-Benzyl benzoate/Diphenyl ether (4:1 vol/vol; BABB-4D) before imaging. Whole E13 embryo heads were recorded with a light sheet fluorescent Ultramicroscope II (Lavision Biotech) utilizing the filters (emission/excitation; Em/Ex) 470/425 nm for tissue autofluorescence and 630/680nm for amplified GFP signal. ARH area was recorded in detail at 4× zoom with a 2-μm plane size. Cell counting was performed with Arrivis Vision4D software (Arrivis) after 3-dimensional (3-D) reconstruction, background removal and cell segmentation.

**Viruses.** Ready-to-transduce lentiviral particles carrying human CUX1 P200 (NM_181552) and RPGRIP1L (NM_015272) were purchased by ORIGENE. AAV1 particles used for stereotaxic injections (AAV1.CMV.PI.eGFP.WPRE.BGH, AAV1.CMV.HI.eGFP-Cre.WPRE.SV40) were purchased from the Penn Vector Core (University of Pennsylvania, Philadelphia, Pennsylvania, USA).

**Cerebellar stereotaxic injections.** Male C57BL/6J mice aged P11 were anesthetized with a combination of ketamine (80 mg/kg) and xylazine (12 mg/kg) (MilliporeSigma, catalog K113), administered i.p. Mice were then placed in a stereotaxic frame adapted for juveniles, and the scalp was clipped of fur and sterilized. A longitudinal incision was made through the scalp, and the skull was leveled. The target site for injection, defined as 2.6 mm caudal to lambda along the midline, was measured stereotaxically, and the
skull above the target was carefully drilled very thin. A glass micropipette loaded in a Drummond Nanoject III device was then filled with AAV1 and positioned over the target site. The pipette was lowered through the soft, thin skull into the cerebellum to a depth of 2.5 mm below the brain surface. Pulses of 20 nl each were injected at 2.5 mm, 2.0 mm, 1.5 mm, 1.0 mm, and 0.5 mm ventral to the brain surface, pausing for 1 minute after each pulse before moving to other sites. Following retraction of the pipette, the incision was closed and mice were permitted to recover from anesthesia prior to being returned to nursing dams.

**ARH and DMH/VMH stereotaxic injections.** Injections of 12-week-old male *Rpgr*−/− mice were performed on a stereotaxic frame using a glass micropipette and an air pressure injector system (76, 77). Each animal was injected at 3 sites bilaterally for a total of 6 injections and a total volume of ~150 nl to ensure complete coverage of the ARH (mediolateral [ML], ±0.3 mm; anteroposterior [AP], –1.6 mm; dorsoventricular [DV], –6 mm) or DMH/VMH (ML, ±0.3; AP, –1.6; DV, –5.8mm). The accuracy of injections was ~50%. Injections that missed the targeted regions did not result in body weight changes. These were excluded from the analysis but provide evidence of the cellular specificity of the phenotypic effects. All surgical procedures were performed using an aseptic technique according to the approved protocols at Columbia University.

**Generation and characterization of iPSCs.** JBST primary fibroblasts were reprogrammed into iPSCs using retroviruses expressing cMYC, OCT4, SOX2, and KLF4 (78). All iPSC lines were cultured on mouse embryonic fibroblast (MEF) cells with human ES medium: 500 ml KO DMEM (Thermo Fisher Scientific, catalog 10829018), 90 ml KO Serum (Thermo Fisher Scientific, catalog A3181502), 6.5 ml GlutaMAX (Thermo Fisher Scientific, catalog 35050061), 6.5 ml NEAA (Thermo Fisher Scientific, catalog 11140050), 6.5 ml penicillin/streptomycin (Thermo Fisher Scientific, catalog 15140122), 0.65 ml β-mercaptoethanol (Thermo Fisher Scientific, catalog 21985023), and 10 ng/ml bFGF (Thermo Fisher Scientific, catalog 13256029). Stable iPSC clones (after 8 passages) were characterized by immunostaining with pluripotency markers SOX2 (1:500; Stemgent, catalog 09-0024), OCT4 (1:500; Stemgent, catalog 09-0023), NANOG (1:500; R&D Systems, catalog AF1997), SSEA4 (1:300; R&D Systems, catalog MAB1435), and TRA-1-60 (1:300; MilliporeSigma, catalog MAB4360), as well as by qPCR analysis to confirm retroviral gene silencing and expression of endogenous pluripotency genes, as described elsewhere (Supplemental Figure 2B) (32, 34). Pluripotency was further confirmed by a teratoma assay: JBST iPSCs (5 million/200µl matrix-gel) were injected s.c. into NSG mice (The Jackson Laboratory, NOD.Cg-Prkdcscid Il2rgtm1Wjl/SzJ, catalog 005557), and after 2–3 months, teratomas were isolated and fixed with 4% paraformaldehyde for >16 hours for immunohistochemical analysis. For karyotyping, control and JBST iPSC lines were cultured in T75 flasks and karyotyped by Cell Line Genetics.

**Neuron differentiation protocols.** Human iPSCs were cultured and maintained in human ESc medium on MEF (111,000 cells/well in 6-well plates; Thermo Fisher Scientific, catalog A34180). Upon confluence, iPSC cells were cultured for 4 days in EB medium (human ESc medium without basic fibroblast growth factor [bFGF] plus dual SMAD inhibitors — A83 (1 µM; Stemgent, catalog 04-0014) and LDN (250 nM; Stemgent, catalog 04-0074). From days 5–8, EB medium was replaced with N2 medium in steps from 75% to 50%, to 25%, and to 0%, while maintaining A83 and LDN at constant concentrations. Neuron progenitors were harvested on day 10 and further expanded for 2–3 passages in N2 medium plus A83/LDN on poly–L-ornithine/Laminin–coated plates. For further neuronal differentiation, cells were cultured on Poly–L-ornithine/Laminin–coated plates (Thermo Fisher Scientific, catalog 08-772-66) at 50,000 cells per 24-well or 4-well plate and at 500,000 cells per 6-well plate. After culturing 3 weeks in N2 medium supplemented with B27 (Thermo Fisher Scientific, catalog 17504404), DAPT (10 µM; Stemgent, catalog 04-0041), and BDNF (20 ng/ml; R&D Systems, catalog 248-BD), neurons were fixed/harvested for IHC or RNA analysis.

H9 and H9-derived isogenic lines were differentiated to ARH neurons following the detailed protocol described in a Methods manuscript by Wang et al. (34).

**CRISPR.** The H9 ES cell line was propagated and subjected to CRISPR as extensively described elsewhere (45). Six isogenic lines per genotype were used for analysis.

**Three-dimensional reconstructions of the PVH/ARH.** We utilized the Zeiss Axio Scan.Z1 (Baymax) automated slide scanner to visualize nuclei by DAPI (Zeiss Filter Set 49), as well as mGFP-positive (FS 38 HE) Pomc neurons and their projections in 60 µm coronal sections spanning the whole PVH and ARH. Images were aligned in Adobe Photoshop; Z-stacks were generated in ImageJ (NIH) and reconstructed in 3-D using Vaa3D (https://github.com/Vaa3D) and CamStudio (www.camstudio.org).

**Statistics.** Data are expressed as means ± SEM unless specified. Statistical analysis was performed using 2-tailed Student’s t test (StatView 5.0, SAS Institute Inc.). Statistical significance of the difference between
3 means was determined by 1-way ANOVA using GraphPad Prism 7. Levels of statistical significance were set at \( P < 0.05 \).

Study approval. Animals were maintained according to Columbia University animal welfare guidelines. All procedures were approved by the Columbia University IACUC. Human subjects provided written informed consent, and human subject–related research was reviewed and approved by the Columbia Stem Cell Committee and the Columbia IRB.

Author contributions
Conceptualization was contributed by LW, CAD, DE, NB, RLL, and GS; methodology was contributed by LW, AJDS, YG, KEB, SEE, CAD, HLA, GMJ, NB, and GS; investigation was contributed by LW, AJDS, YG, KEB, SEE, RT, JML, XL, RR, MP, JYL, JT, MCDR, and GS; resources were contributed by CAD, HLA, JCB, DE, RC, NB, RLL, and GS; writing the original draft was contributed by GS; writing, reviewing, and editing the draft were contributed by LW, AJDS, YG, KEB, CAD, JYL, NB, RLL, and GS; and supervision was contributed by CAD, HLA, JCB, DE, NB, RLL, and GS.

Acknowledgments
We thank Joel Elmquist (UT Southwestern, Dallas, Texas, USA) and Daniel Doherty (University of Washington, Seattle, Washington, USA) for providing the POMC-CreER mice and JBST fibroblast lines, respectively. Also, we thank Lisa Cole Burnett (Columbia University) for microscopy. Work was partially funded by a Russell Berrie Foundation Neuroscience of Obesity Initiative pilot grant, RO1 DK52431-23, and the MBMG Core of the NYNORC (SP30DK026687), the New York Stem Cell Foundation, the Rudin Foundation, the Russell Berrie Foundation Program in Cellular Therapies, and the Diabetes (P30 DK63608-12) and Obesity Research (P30 DK26687-33) Centers of Columbia University. Yossef Goffer was supported by NIDDK F30DK108564. Nicolas F. Berbari was supported by RO1 DK114008 and CDMD NIH/NIDDK P30 DK097512. Alain J. De Solis was supported by MSCA-IF-2016 (752319, ARC-Neurohet).

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