

High-Throughput Screening of Mouse Gene Knockouts Identifies Established and Novel High Body Fat Phenotypes

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Purpose: Obesity is a major public health problem. Understanding which genes contribute to obesity may better predict individual risk and allow development of new therapies. Because obesity of a mouse gene knockout (KO) line predicts an association of the orthologous human gene with obesity, we reviewed data from the Lexicon Genome5000TM high throughput phenotypic screen (HTS) of mouse gene KOs to identify KO lines with high body fat.

Materials and Methods: KO lines were generated using homologous recombination or gene trapping technologies. HTS body composition analyses were performed on adult wild-type and homozygous KO littermate mice from 3758 druggable mouse genes having a human ortholog. Body composition was measured by either DXA or QMR on chow-fed cohorts from all 3758 KO lines and was measured by QMR on independent high fat diet-fed cohorts from 2488 of these KO lines. Where possible, comparisons were made to HTS data from the International Mouse Phenotyping Consortium (IMPC).

Results: Body fat data are presented for 75 KO lines. Of 46 KO lines where independent external published and/or IMPC KO lines are reported as obese, 43 had increased body fat. For the remaining 29 novel high body fat KO lines, *Ksr2* and *G2e3* are supported by data from additional independent KO cohorts, 6 (*Asnsd1*, *Srpk2*, *Dpp8*, *Cxxc4*, *Tenm3* and *Kiss1*) are supported by data from additional internal cohorts, and the remaining 21 including *Tle4*, *Ak5*, *Ntm*, *Tusc3*, *Ankk1*, *Mfap3l*, *Prok2* and *Prokr2* were studied with HTS cohorts only.

Conclusion: These data support the finding of high body fat in 43 independent external published and/or IMPC KO lines. A novel obese phenotype was identified in 29 additional KO lines, with 27 still lacking the external confirmation now provided for *Ksr2* and *G2e3* KO mice. Undoubtedly, many mammalian obesity genes remain to be identified and characterized.

Keywords: obesity, druggable, homologous recombination, gene trapping

Introduction

The obesity pandemic is a major public health problem. The global prevalence of obesity has increased steadily since 1980, and by 2015 108 million children and 604 million adults worldwide were obese.^{1,2} In the United States, the prevalence of obesity in adults increased from 33.7% in 2007–2008 to 42.4% in 2017–2018.³ These obesity estimates are based on body-mass index (BMI) data, calculated as weight/height² (kg/meter²), with obesity defined as BMI \geq 30. High BMI is a risk factor for many chronic diseases; in 2015, high BMI was linked globally to 4

million deaths and 120 million disability-adjusted life years.^{2,4} Clearly, there is a need for new interventions that effectively decrease the amount of body fat.

Human obesity, as represented by the surrogate measure BMI, has a genetic component that explains ~30% of the variability.⁵⁻⁹ The value in identifying genes responsible for this heritability is the potential to predict individual risk for future obesity and the potential to identify pathways and targets for therapeutic intervention. The first obesity genes were discovered by positional cloning studies using the obese mouse lines *ob/ob* and *db/db*, and by studying the agouti yellow obese mouse; these efforts identified leptin (LEP), leptin receptor (LEPR) and melanocortin 4 receptor (MC4R) as integral proteins regulating body fat in mice and humans. Investigating the role of these proteins in neural pathways controlling adiposity resulted in identification of a few additional genes that help to regulate mammalian body fat stores.¹⁰⁻¹⁴ To date, identifying these obesity genes has not led to therapies capable of reversing the obesity pandemic.

The inability of early linkage studies to identify loci linked to common obesity suggested correctly that numerous genetic variants make small individual contributions; currently, >250 GWAS loci are identified as minor contributors to BMI variance.^{8,15-18} Attempts to integrate these GWAS loci into a single genome-wide polygenic score that predicts individual obesity risk had limited success.¹⁹ Also, these GWAS loci rarely pinpoint novel therapeutic targets because most reside in introns as clusters of noncoding variants which likely regulate expression, not function, of a causal protein encoded by a gene which may be located far from the locus itself.¹⁶ A good example is the locus located in the first intron of *FTO* (fat mass and obesity associated). A role in obesity for the demethylase enzyme encoded by *FTO* is supported by loss of function mutations in mice, but not in humans,^{20,21} and recent studies link these *FTO* variants to altered expression of the nearby *RPGRIP1L* and distant *IRX3* and *IRX5* genes, with loss of function mutations of *Irx3* and *Rpgrip1l* genes altering energy balance in mice.²⁰⁻²² Of these 4 candidate genes, only *FTO* encodes a druggable protein by conventional standards, where druggable proteins are either secreted, accessible in vivo to a circulating antibody, or members of a protein family successfully targeted by small molecules.²³⁻²⁵ After a decade of research, the *FTO* locus has not yielded a mature drug discovery program. Most recently, an exome-wide search for low frequency and rare protein-altering variants (allele

frequency < 5%) associated with BMI identified 8 novel obesity genes.¹⁶ These variants contributed little to BMI variation due to their rarity. However, this exome-sequencing approach has value over previous GWAS approaches because the genes responsible are identified.

Inactivating mutations in mouse genes often identify genes that regulate human body fat stores,²⁶ clarify the physiologic role of GWAS-associated human genes,²⁷ and mimic phenotypes resulting from human gene mutations.²⁸ Also, mice with knockout (KO) mutations that inactivate the targets of drugs usually exhibit phenotypes consistent with the effects of those drugs in humans; this correlation between effects of genetic manipulation in mice and pharmacologic manipulation in humans suggests conservation of gene function.^{29,30} This correlation extends to body fat regulation; of 21 obesity gene targets reviewed, most showed a correspondence between KO phenotype and drug effect in rodents, and, when data existed, drug effect in humans.^{31,32} Together, these data suggest that coordinated KO of all mouse genes followed by phenotyping of viable KO lines for body fat content will identify genes playing a previously unrecognized role in human obesity.

Because mouse global KO phenotypes model drug effects, Lexicon Pharmaceuticals pursued, between 2000 and 2008, the high-throughput Genome5000TM program designed to KO and phenotype the druggable genome and reveal novel drug targets.^{29,33-37} We have reported the skeletal phenotypes identified by the primary high-throughput screen (HTS).³⁷ The HTS also included assessment of body fat in two independent mouse cohorts fed either high fat diet (HFD) or chow.³⁵ In a preliminary report, we showed that the HTS correctly identified body fat phenotypes in 13 benchmark KO lines.³⁶ The present report extends these observations and summarizes our experience identifying established and novel high body fat phenotypes in our screen of the druggable genome.

Methods

Mouse Care and Study

All animal studies were performed in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. Protocols for all studies were approved by the Lexicon Institutional Animal Care and Use Committee (OLAW Assurance Number, A4152-01; AAALAC International Accreditation Number, 001025). General methods for mouse care are published.^{36,38,39} Briefly, WT

and KO mice were housed together, with a maximum of 5 mice/cage, in a temperature-controlled environment on a fixed 12-hour light/12-hour dark cycle and with free access to food and water. Unless stated otherwise, mice received standard rodent chow 9F 5020 (Purina, St. Louis, MO) as our chow diet, D12450B (Research Diets, New Brunswick, NJ) containing 10% of kcals from fat as our low-fat diet (LFD), or D12451 (Research Diets) containing 45% of kcals from fat as our high-fat diet (HFD). In a few studies, mice were fed a high-fat diet containing 60% kcals from fat (60% HFD; D12492, Research Diets). Pair-feeding studies were performed as described previously.⁴⁰

Generation of Knockout Mice

Our approach to KO and phenotype mouse orthologs of the estimated 5000 potential drug targets in the human genome (Genome 5000TM project) is published.^{29,34–37,41–43} Briefly, Lexicon generated KO lines for the Genome 5000TM project by manipulating the mouse genome using either gene trapping (OmniBank[®] I) or homologous recombination technologies. Gene trapping mutagenesis in embryonic stem cells involves inserting DNA elements randomly into endogenous genes leading to transcriptional disruption and introduction of sequence tags. Oligonucleotide primers complementary to the gene-trap vector precisely localize the vector insertion site within the exon-intron organization of the gene, and gene disruption is confirmed using RT-PCR performed with oligonucleotide primers complementary to exons flanking the vector insertion site.³⁷ This method permitted selection of druggable genes for KO generation. Because many factors place an upper limit on the percentage of genes that are efficiently trapped, gene KO by homologous recombination, which allows targeting of selected genes including those not targeted by gene trapping, was also employed. All mutations were generated in strain 129SvEvBrd-derived ES cells. Resulting chimeric mice were bred to C57BL/6J albino mice to generate F1 heterozygous (HET) progeny. These mice were intercrossed to generate F2 wild-type (WT), HET and KO (homozygous mutant) offspring, which were studied in our high-throughput primary phenotypic screen. All mice were on a C57BL/6J X 129SvEvBrd hybrid background. Genotyping was performed on tail DNA as described.⁴⁴

[Supplementary Table 1](#) lists official names and gene symbols for every gene with a KO line discussed here. The strategies for generating many of these KO lines are published, including *Asnsd1*;⁴⁵ *Htr2c*, *Brs3*, *Mc4r*, *Mc3r* and

Tsn;³⁶ *Ffar4* or *Gpr120*;⁴⁶ *G2e3*;⁴⁷ *Hdac5*;⁴⁸ *Ksr2*;^{36,49} *Prlhr*;⁵⁰ *Slc6a4* or *Sert*;⁵¹ and *Kiss1*, *Kiss1r*, and *Sost*.³⁷ The *Mc3r* and *Mc4r* KO lines were used to generate *Mc3r/Mc4r* double KO (DKO) mice. [Supplementary Table 2](#) presents homologous recombination strategies for KO lines of *Adamts4*, *Adamts18*, *Adm2*, *Ak5*, *Ankk1*, *Apln*, *Aqp7*, *AU040320*, *Ccn5*, *Cxxc4*, *Dgkg*, *Dpp8*, *Gpr45*, *Gpr61*, *Gpx7*, *Itih1*, *Lrrn2*, *Lrrtm1*, *Mfap3l*, *Ncs1*, *Npvf*, *Ntm*, *Oprm1*, *P2rx6*, *Pnpla2*, *Prok2*, *Prokr2*, *Ptpn*, *Pyy*, *Resp18*, *Retn*, *Retsat*, *Scg3*, *Sik2*, *Tnfrsf13b*, *Usp13*, *Usp38* and *Wnt8b*. [Supplementary Table 3](#) describes the location of gene trap insertion vectors for KOs of *Adamts4*, *Adcy3*, *Aoc3*, *Ddah1*, *Enox1*, *Glr2*, *Hdac6*, *Herc1*, *Igfbp2*, *Igdcc4*, *Kdm3a*, *Ncoa1*, *Nr4a1*, *Pecr*, *Prdx6*, *Prmt7*, *Ptp4a1*, *Rgs10*, *Scg3*, *Srp2*, *St3gal2*, *Tenm3*, *Tle4*, and *Tusc3*.

Primary High-Throughput Screen (HTS)

Our primary HTS consisted of a battery of tests performed on F2 mice, including audiology, behavior, hematology, serum chemistry, bone densitometry, histopathology, cardiology, metabolism, immunology, oncology and ophthalmology assays.^{35,37} Briefly, 2 cohorts of mice were studied. A cohort fed chow diet from weaning usually consisted of 4 WT, 4 HET and 8 KO mice, half male and half female, for each project. A cohort fed HFD from weaning consisted of up to 8 WT and 8 KO male mice for each project.

As part of this HTS, body composition was analyzed on both mouse cohorts. A cohort of 14-week old mice fed chow from weaning was screened, usually by dual-energy X-ray absorptiometry (DXA) using a GE/Lunar PIXImus scanner (GE Medical Systems, Madison, WI) but in a few instances by quantitative magnetic resonance (QMR) technology using a Bruker Minispec QMR Analyzer (ECHO Medical Systems, Houston, TX).³⁶ Briefly, for each KO line, the mean KO % body fat/mean wildtype (WT) littermate % body fat ratio was calculated for both male and female mice, and then these male and female values were averaged to yield a normalized % body fat (n%BF) value. For most KO lines, 4 male KO, 2 male WT, 4 female KO and 2 female WT mice were studied. KO lines with fewer than 4 KO or 3 WT mice were excluded, except for X-linked lines where only male data were used, usually from 8 KO and 2 WT mice. A separate cohort of 11-week old male mice fed HFD from weaning was screened by QMR as described previously;⁴⁹ all lines with between 3 and 8 KO mice and between 3 and 8 WT littermate

controls were included in the analysis. The n%BF for each line was calculated as described for the chow-fed cohort. Also, for some KO lines, normalized body fat grams (nBFg) was calculated from BFg data and normalized lean body mass (nLBM) was calculated from LBM data, as described for n%BF. Historical mean and SD values for %BF and BFg were calculated using data from ~14500 WT mice studied in the chow-fed cohorts from 3758 independent KO lines and from ~16000 WT mice studied in the HFD-fed cohorts from 2475 independent KO lines.

Studies Using Additional Cohorts

Selected KO lines with high body fat identified in the primary chow and HFD screens were studied using additional mouse cohorts to determine if the HTS high body fat phenotype was reproducible. Additional KO lines were selected for further study based on published reports linking that KO to obesity. These body composition measurements were made using QMR or DXA technologies as described above. In addition to body composition, additional phenotyping assays were performed; methods for these assays are described below.

Blood Sample Analysis

Unless stated otherwise, blood was obtained from fed mice by retro-orbital bleed and serum assayed for glucose and total cholesterol by Cobas Integra 400 analyzer (Roche Diagnostics, Indianapolis, IN).^{35,39} Serum leptin levels were measured using an ELISA kit (Mouse Leptin, Crystal Chem, Downers Grove, IL).

VO₂ and Physical Activity

VO₂ and ambulatory activity were measured in OxyMax chambers (OxyMax, Columbus Instruments, Columbus, OH) as described previously.⁴⁹ Gross motor activity was measured using the ER-4000 physiological measurement system (Mini Mitter, Bend, OR, USA). Three days after E-mitter transponders were surgically implanted into their peritoneal cavity, mice were placed in individual cages within range of an ER-4000 receiver, which measures activity by sensing the strength of the signal received from the E-mitter. Activity data for each mouse were collected in 10 min intervals for a total of 6 days (144 hours) using the VitalView Data Acquisition System (Mini Mitter); gross motor activity data were normalized such that mean WT activity = 100%.

Glucose Tolerance Tests

Glucose tolerance tests (GTTs) were performed on conscious, unanesthetized mice. After an overnight fast, mice were bled at baseline and then received 2 g/kg glucose by oral gavage (OGTT) or by intraperitoneal injection (IPGTT). Glucose levels from each mouse, obtained on whole blood samples at 0 (baseline), 30, 60, 90 and/or 120 minutes using an ACCU-CHEK Aviva glucometer (Roche, Indianapolis, IN) were used to calculate glucose area under the curve (AUC) values. Two additional serum aliquots obtained at 0 (baseline) and 30 minutes were used to measure insulin levels (Ultra-Sensitive Rat Insulin ELISA Kit, Cat. 90060; Crystal Chem).

Fertility Testing

For each project, 1–3 male and 2–4 female KO mice were examined. Each male KO mouse was bred to 2 WT females for 14 days. Each female KO mouse was bred individually with up to 3 WT males for 60 days. Pregnant females (KO and WT) were monitored daily for births; pups were counted at birth and at postnatal day 4 and were monitored daily for stomach milk content and for growth. Special care was taken not to disturb the litter environment.

Systolic Blood Pressure

Systolic blood pressure (SBP) was measured on conscious 10 to 13 week-old mice using a tail-cuff system (Visitech Systems, Apex, NC, USA). SBP was measured 10 times daily for 4 consecutive days, and the SBP value reported was the mean of the 40 SBP readings.⁵²

Histopathology

Methods for general histopathology are published.⁴⁹

External Public Database Resources

Human GWAS data identifying clusters of BMI, waist-hip ratio adjusted BMI, body fat percentage, weight and T2D single nucleotide polymorphism variants located within 1.2 MBP 5' or 3' of selected genes were obtained at <https://hugeamp.org/> which links to the Common Metabolic Diseases Knowledge Portal; the 1.2 MBP distance was chosen because that is the distance at which *FTO* intronic variants influence *IRX5* expression.²² For each gene, the distance between the cluster variant closest to the gene and the gene exon closest to that variant was calculated. Human gene expression data were obtained using the GTEx Portal at <https://gtexportal.org/home/>

[GTEx Analysis Release V8 (dbGaP Accession phs000424.v8.p2)]. HTS data for body weight (BW), body fat, fertility and/or total cholesterol levels from selected mutant mouse gene KO lines were obtained using the International Mouse Phenotyping Consortium (IMPC) database at <https://www.mousephenotype.org> (Data Release Version 14). The IMPC HTS typically examined 7 male and 7 female mutant and WT mice with body composition measured by DXA.⁵³ Catalog designations and additional information on Lexicon KO models available at Taconic Biosciences were found at <https://www.taconic.com/find-your-model/>.

Statistics

Data are presented as mean \pm SD. Comparisons between two groups were analyzed by unpaired Student's *t* test; when variances between the two groups were significantly different, the data were analyzed using the nonparametric Mann–Whitney test. Comparisons among three groups were analyzed by one-way ANOVA with Tukey's test used to correct for multiple comparisons. Two-way ANOVA was performed on body fat and LBM data from *Mc3r* KO, *Mc4r* KO, *Mc3r/Mc4r* DKO and WT littermate mice. Viability of KO lines was determined by chi-square testing of Mendelian ratios obtained at weaning. All statistical tests were performed using PRISM 4.03 (GraphPad Software, Inc., La Jolla, CA, USA). Differences were considered statistically significant when $P < 0.05$.

Results

We obtained HTS body composition data on 3758 independent mouse gene KO lines with viable adult homozygous KO mice. Body fat was measured on chow-fed cohorts of WT and KO littermate mice from all 3758 KOs; 3650 KO lines were studied by DXA and 108 KOs were studied by QMR. For 2488 of these KO lines, body fat was also measured by QMR on an independent cohort of high fat diet-fed WT and KO littermate mice. HET mice, along with their WT littermates, were also examined in our HTS for 755 KO lines exhibiting homozygous lethality. For all but three genes with viable KO mice, the HET data were not analyzed because there were only 4 HET mice/KO line. For *Adcy3*, which is one of 90 KO lines showing reduced viability of homozygous KO mice prior to 10 weeks of age, data were pooled from 2 KO and 8 HET littermates. For *Scg3*, which had 2 independent KO lines, and *Adamts4*, which had 3 independent KO lines, data were pooled from 12 HET and 16 KO littermates, and

from 12 HET and 24 KO littermates, respectively. For each of these three genes where data were available for at least 8 HET mice, HET and KO littermate mice showed a similar increase in body fat.

Obese KO Lines Studied with Both HTS and Additional Cohorts

Figure 1 shows HTS normalized percent body fat (n%BF) values for each of these KO lines within the distribution of values for all KO lines fed chow diet from weaning. Table 1 presents pooled n%BF data that combined, for each KO line, the HTS cohort data shown in Figure 1 with data from all additional cohorts. These obese KO lines studied with both HTS and additional cohorts are categorized here as follows: 1) Benchmarks, consisting of the 5 KO lines with increased body fat reported in two external publications that we used previously to validate our HTS,³⁶ a crucial step which showed that, despite examining a small number of mice/KO line, our HTS was unlikely to report many false negatives in which high body fat was missed in specific gene KO lines; 2) KO lines first published by external groups that did not originally serve as benchmarks; and 3) Novel KO lines which consist of KO lines, either published or unpublished, that were first identified at Lexicon. For all normalized data, the mean WT value for each cohort is assigned a value of 100%. Table 2 presents human expression and associated GWAS BMI cluster data for selected non-benchmark KO lines presented in Table 1. Human expression data focus on CNS, hypothalamic and adipose tissues to provide insight into how the targeted gene might regulate body fat, because most obesity genes are expressed in the CNS and likely regulate food intake centrally through hypothalamic relays, while a few may have a direct effect on adipogenesis and/or fat storage by adipocytes.^{8,54} Data on human GWAS BMI clusters located close to the human orthologs of targeted mouse genes are presented to suggest that these human orthologs may be responsible for the GWAS BMI signals.

Benchmark KO Lines

We originally showed, using 5 benchmark KO lines with established obesity (*Brs3*,^{55,56} *Tsn*,^{57,58} *Mc4r*,^{13,59,60} *Mc3r*^{60–62} and *Htr2c*^{63,64}) that our HTS correctly predicted the amount of body fat present in additional cohorts.³⁶ Our data confirm published data showing that KO models for *Brs3*, *Tsn*, *Mc4r*, and *Mc3r* are obese. Our data also confirm that *Tsn* KO mice have low LBM, with nLBM of KO

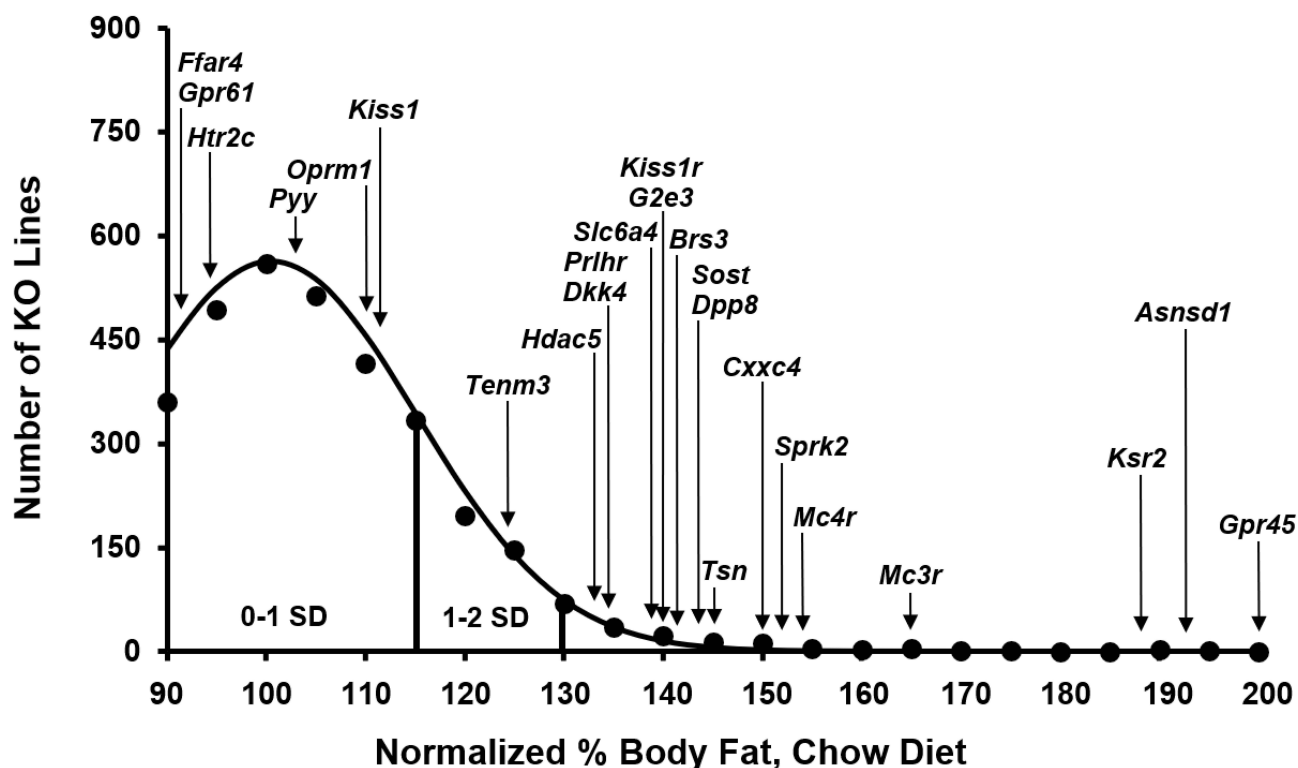


Figure 1 High-throughput screen (HTS) normalized %body fat (n%BF) values for individual well-studied KO lines within the distribution of HTS n%BF values for all 3650 individual KO lines maintained on chow diet from weaning. Body composition analyses performed by DXA on 14-week-old mice were used to calculate n%BF for the cohort from each individual KO line. Solid points indicate actual numbers of KO lines within that mean \pm 2.5% value of n%BF. Curved line shows the calculated curve. The range for 1 and 2 SD from the population mean is indicated by lines located below the curve, and the mean n%BF value for the HTS cohort from each individual well-studied KO line is indicated by arrows above the curve. Some of these KO lines also had HTS data generated on an independent HFD-fed cohort; these KO lines and their n%BF data from the HFD-fed HTS cohort were: *Pyy* (101%), *Sprk2* (106%), *Dkk4* (107%), *Htr2c* (108%), *Tenm3* (110%), *Prlhr* (111%), *Oprm1* (123%), *Hdac5* (132%), *Dpp8* (134%), *Mc4r* (159%), *Kiss1* (161%), *G2e3* (164%), *Asnsd1* (167%) and *Ksr2* (189%).

mice = 89 ± 9 (n=21) vs WT mice = 100 ± 5 (n=13), $P < 0.001$; however, in contrast to published data, our *Tsn* KO mice had impaired glucose tolerance with OGTT glucose AUC of KO mice = 33227 ± 12131 (n=8) vs WT mice = 22978 ± 4187 mg*min/dL (n=9), $P < 0.05$ by Mann Whitney test. In addition, analysis of our cohort of *Mc3r/Mc4r* DKO mice showed that the increased BFG of DKO mice is comparable to the combined increase observed in *Mc3r* and *Mc4r* KO mice, suggesting independent pathways of fat accumulation, and that only *Mc4r* KO mice have increased LBM (Supplementary Table 4), all confirming published data.^{60,62} In contrast, our *Htr2c* KO mice did not show significantly increased body fat either in the HTS or in additional cohorts³⁶ despite the obesity reported in 2 independent *Htr2c* KO lines and the ability of the HTR2C inhibitor lorcaserin to both lower body fat in obese humans and lower food consumption in *Htr2c* WT, but not KO, mice.^{63–66} A review of HTS data for these benchmark KO lines found that the number of KO mice with %BF or BFG > 2 SD above the historic

mean, compiled with data from ~14500 WT mice in the chow fed HTS and ~16000 WT mice in the HFD HTS, was 15/16 for *Mc4r*, 3/8 for *Mc3r*, *Brs3* and *Tsn*, and 1/16 for *Htr2c*; in contrast, the number of WT mice was 1/11 for *Mc4r*, 0/4 for *Mc3r*, *Brs3* and *Tsn*, and 1/10 for *Htr2c*. This suggests that KO lines with marked obesity have many KO mice that are quite obese.

Additional Externally Published KO Lines Not Used as Benchmarks

GPR45 is an orphan GPCR with constitutive activity. External *Gpr45* KO mice had increased body fat at weaning and developed progressive obesity due mainly to decreased energy expenditure, a conclusion based on studies performed during the development of obesity which showed that KO mice had comparable food intake, decreased energy expenditure and decreased physical activity relative to WT mice; these KO mice were also glucose intolerant with high plasma insulin and leptin levels, and displayed reduced hypothalamic proopiomelanocortin (POMC) expression suggesting a link to the

Table I Normalized % Body Fat Data for KO Lines with Both HTS and Secondary Screen Cohorts

Normalized % Body Fat, Mean \pm SD (n)						
KO Line	KO	WT	P value	Sex	Age (Weeks)	Diet
Benchmark KO lines						
<i>Mc3r</i>	176 \pm 36 (43)	100 \pm 26 (35)	<0.001	M and F	5–14	Chow and HFD
<i>Mc4r</i>	164 \pm 41 (104)	100 \pm 21 (107)	<0.001 [†]	M and F	7–21	Chow and HFD
<i>Tsn</i>	160 \pm 43 (21)	100 \pm 17 (13)	<0.001 [†]	M and F	14–20	Chow
<i>Brs3</i>	148 \pm 42 (29)	100 \pm 34 (13)	<0.001	M (hemi)	14–19	Chow
<i>Htr2c</i>	105 \pm 25 (51)	100 \pm 20 (48)	0.268	M (hemi)	11–35	Chow and HFD
Additional published high body fat KO lines						
<i>Gpr45</i>	205 \pm 59 (118)	100 \pm 23 (93)	<0.001 [†]	M and F	7–38	Chow and HFD
<i>Kiss1r</i>	140 \pm 32 (18)	100 \pm 13 (13)	<0.001 [†]	M and F	14–16	Chow
<i>Gpr61</i>	125 \pm 31 (68)	100 \pm 25 (57)	<0.001	M and F	14–21	Chow and HFD
<i>Hdac5</i>	125 \pm 30 (35)	100 \pm 20 (25)	<0.001	M and F	11–25	Chow and HFD
<i>Prlhr</i>	120 \pm 36 (54)	100 \pm 22 (33)	0.002 [†]	M and F	11–24	Chow and HFD
<i>Sost</i>	117 \pm 26 (52)	100 \pm 24 (56)	<0.001	M and F	14–30	Chow
<i>Slc6a4</i>	117 \pm 29 (49)	100 \pm 29 (48)	0.005	M and F	9–50	Chow
<i>Retn</i>	117 \pm 20 (27)	100 \pm 20 (31)	0.002	M and F	11–24	Chow and HFD
<i>Oprm1</i>	116 \pm 15 (26)	100 \pm 23 (29)	0.008	M and F	11–16	Chow and HFD
<i>Pyy</i>	102 \pm 24 (25)	100 \pm 21 (21)	0.765	M and F	11–37	Chow and HFD
<i>Ffar4</i>	99 \pm 17 (38)	100 \pm 12 (23)	0.843	M and F	11–14	Chow and HFD
Novel high body fat KO lines						
<i>Ksr2</i>	235 \pm 46 (127)	100 \pm 21 (153)	<0.001 [†]	M and F	5–14	Chow and HFD
<i>Asnsd1</i>	180 \pm 44 (39)	100 \pm 24 (34)	<0.001 [†]	M and F	11–42	Chow and HFD
<i>G2e3^a</i>	148 \pm 32 (32)	100 \pm 28 (31)	<0.001	M and F	11–25	Chow and HFD
<i>Srpk2</i>	141 \pm 38 (41)	100 \pm 26 (48)	<0.001 [†]	M and F	11–22	Chow and HFD
<i>Dpp8</i>	133 \pm 23 (39)	100 \pm 17 (32)	<0.001	M and F	11–28	Chow and HFD
<i>Cxxc4</i>	131 \pm 39 (39)	100 \pm 24 (30)	<0.001 [†]	M and F	14–23	Chow
<i>Kiss1</i>	127 \pm 31 (27)	100 \pm 30 (24)	0.003	M and F	11–27	Chow and HFD
<i>Tenm3</i>	121 \pm 31 (29)	100 \pm 25 (34)	0.004	M and F	11–15	Chow and HFD

Notes: KO mice in each group presented by decreasing % body fat; [†] Statistical analysis by Mann–Whitney test. ^aData published previously in reference 47.

Abbreviations: SD, standard deviation; n, number of mice; KO, knockout; WT, wild-type; M, male; F, female; hemi, hemizygous; HFD, high fat diet.

melanocortin pathway.⁶⁷ Our *Gpr45* KO mice, first reported as the LG747 KO line,⁶⁸ also had increased body fat at weaning and developed progressive obesity that was still present at 87 weeks of age ([Supplementary Table 5](#)). Although we did not observe differences in VO₂ or physical activity between KO and WT littermates ([Supplementary Table 6](#)), a pair feeding study initiated at weaning showed clearly that obesity developed in male and female *Gpr45* KO mice despite food intake that was comparable to that of their WT littermates ([Supplementary Figure 1](#)). Although we did not observe impaired glucose tolerance in our KO mice, we found increased insulin levels associated with severe insulin resistance, and increased serum levels of leptin, triglycerides and total cholesterol ([Supplementary Table 7](#)).

KISS1 is a neuropeptide that activates the hypothalamic-pituitary-gonadal axis by binding the hypothalamic G protein coupled receptor (GPCR) KISS1R; inactivating mutations in either gene result in delayed or absent puberty in mice and humans.⁶⁹ Our *Kiss1* and *Kiss1r* KO mice exhibited male and female hypogonadism and showed decreased bone mass consistent with hypogonadism.³⁷ Published studies also found increased body fat in *Kiss1r* KO mice, more striking in females, and decreased LBM, more striking in males; these findings suggest a metabolic role for these genes.^{69,70} Our data confirm these findings for *Kiss1r* KO mice and show the same pattern for *Kiss1* KO mice except that both male and female *Kiss1* KO mice had increased body fat ([Supplementary Table 8](#)).

Table 2 Human Expression and GWAS Data on Selected Non-Benchmark Obese KO Lines with HTS and Secondary Screen Cohorts

		Major GWAS BMI Clusters Within 1.2 MBP ^b		
KO Line	Human Expression ^a	Lead Variant	P value	Cluster Distance
Additional KO lines with published high body fat				
<i>Gpr45</i>	Mainly CNS, highest in Hyp	2:105976344:A:G	<5e-8	0.06 MBP 3'
<i>Kiss1r</i>	Mainly CNS, mainly Hyp; Pit	19:1956035:G:A	<e-14	0.89 MBP 3'
<i>Gpr61</i>	Mainly CNS, incl Hyp; Pit	1:110154688:T:C	<e-38	Overlapping
<i>Hdac5</i>	Wide; CNS, incl Hyp; Ad	17:42193185:G:C	<e-11	Overlapping
<i>Prlhr</i>	Mainly Pit, Hyp	10:120465796:C:G	<e-10	0.07 MBP 5'
<i>Sost</i>	Mainly Osteocytes, Lung; not CNS	17:42193185:G:C	<e-11	0.33 MBP 5'
<i>Slc6a4</i>	Mainly lung, GI	17:28410277:A:G	<e-11	Overlapping
<i>Oprm1</i>	Mainly CNS, incl Hyp	6:154309808:C:T	<e-14	Overlapping
Novel high body fat KO lines				
<i>Ksr2</i>	Mainly CNS, incl Hyp	12:118409274:G:A	<e-10	Overlapping
<i>Asnsd1</i>	Wide; CNS, incl Hyp; Ad, SkM	None		
<i>G2e3</i>	Wide; CNS, incl Hyp; Ad	14:30495719:C:T	<e-15	0.49 MBP 5'
<i>Srpk2</i>	Wide; CNS, incl Hyp; Ad	7:104756355:T:C*	<e-15	Overlapping
<i>Dpp8</i>	Wide; CNS, incl Hyp; Ad	15:65916527:A:T	<5e-8	Overlapping
<i>Cxxc4</i>	Wide; CNS highest, mainly Hyp; Ad	4:106160133:G:A*	<e-11	0.65 MBP 5'
<i>Kiss1</i>	Hyp only	1:203491392:T:G	<e-11	0.67 MBP 3'
		1:204414769:T:C*	<5e-8	0.25 MBP 5'
<i>Tenm3</i>	Wide; CNS, incl Hyp; Ad	None		

Notes: KO mice in each group presented by decreasing % body fat. ^aData obtained from <https://gtexportal.org/home/>. ^bData obtained from <https://hugeamp.org/> on 8/17/2021. *Waist-hip ratio adjusted BMI.

Abbreviations: MBP, million base pairs; incl, including; CNS, central nervous syndrome; Hyp, hypothalamus; Ad, adipose; Pit, pituitary; GI, gastrointestinal tract; SkM, skeletal muscle.

GPR61 is an orphan GPCR with constitutive activity. *Gpr61* KO mice were reported as hyperphagic and obese with decreased hypothalamic *Pomc* expression that potentially links this GPR to the melanocortin pathway.⁷¹ Although *Gpr61* KO mice were not obese in our HTS (Figure 1), sequence similarity of GPR61 to GPCRs activated by biogenic amines suggested druggability. This led us to study additional cohorts where we found that our *Gpr61* KO mice were obese, with a 25% increase in n%BF (Table 1), a 45% increase in nBFg (KO=145 ± 56 vs WT=100 ± 35, $P<0.001$ by Mann-Whitney test) and a 4% increase in nLBM (KO=104 ± 14 vs WT=100 ± 8, $P<0.001$ by Mann Whitney test).

HDAC5 is an enzyme that influences hypothalamic leptin signaling, in part by STAT3 deacetylation.⁷² Our data confirmed that *Hdac5* KO mice are obese; our KO mice had a 25% increase in n%BF (Table 1), a 19% increase in nBFg (KO=119 ± 41 vs WT=100 ± 29, $P<0.05$) and a 12% decrease in nLBM (KO=88 ± 12 vs WT=100 ± 7, $P<0.001$).

PRLHR is a GPCR that confers the ability of prolactin-releasing peptide (PrRP) to decrease food intake and increase energy expenditure, a role supported by the obesity observed in two independent *Prlhr* KO models^{50,73} and on the anorectic effects of PrRP when delivered to rats and *Prlhr* WT, but not KO, mice.^{74,75} Our studies confirm obesity in *Prlhr* KO mice; the KO mice showed a 20% increase in n%BF (Table 1) and a 35% increase in nBFg (KO=135 ± 58 vs WT=100 ± 32, $P<0.01$).

The serotonin reuptake transporter SLC6A4 regulates serotonin action and is a target of antidepressants that may affect food intake and BW. The obesity observed in KO mice by two groups studying the same *Slc6a4* KO mouse model^{76,77} was confirmed in KO mice from our independent *Slc6a4* KO line (Table 1).

RETN, a protein secreted by adipocytes in mice and macrophages in humans, impairs insulin action in rodents.^{78,79} *Retn* KO mice did not have increased BW but *ob/ob* mice lacking RETN were obese compared to

ob/ob mice,⁷⁸ as were transgenic mice expressing a dominant inhibitory form of RETN.⁷⁹ Our *Retn* KO mice were modestly obese, consistent with the published data.

The *SOST* gene encodes sclerostin, an inhibitor of WNT signaling secreted mainly by osteocytes.⁸⁰ In humans and mice, inactivating *Sost* mutations increase bone mineral density and strength.^{37,81} Recently, *Sost* KO mice and mice receiving sclerostin-neutralizing antibodies were found to have low body fat, while mice overexpressing sclerostin were obese, apparently due to altered WNT signaling in adipocytes.⁸² However, older work reported obesity in *Sost* KO mice,⁸³ consistent with the finding that mice with an osteocyte-specific increase in sclerostin release had lower body fat and increased beiging of adipocytes;⁸⁰ our data support these last two studies, with *Sost* KO mice showing modest increases of 17% in n%BF (Table 1) and 16% in nBFg (KO=116 ± 38 vs WT=100 ± 36, $P<0.05$ by Mann–Whitney test). This link between sclerostin and obesity is intriguing given recent evidence that a leptin-independent BW homeostat is dependent on osteocytes.⁸⁴

The opioid system, particularly OPRM1, participates in neural reward processes leading to addictive behavior; many neural structures involved in addictive behavior are also involved in food reward, and opioid receptor antagonists inhibit intake of both addictive drugs and palatable diet.⁸⁵ Consistent with these findings, %BF was lower in *Oprm1* KO mice maintained on highly palatable HFDs;^{86,87} surprisingly, however, some but not all past studies found increased BW and/or %BF in *Oprm1* KO mice fed less palatable chow diets.^{86–88} *Oprm1* KO mice reported here had increased %BF (Table 1) and nBFg (KO=125 ± 20 vs WT=100 ± 29, $P<0.001$), with a similar increase in body fat for KO mice fed chow or HFD. These data suggest that *Oprm1* KO mice may develop modest obesity when fed certain diets.

PYY is a gastrointestinal peptide that induces central satiety. In some *Pyy* KO mouse studies, body fat was increased in male KO mice fed chow⁸⁹ or HFD⁹⁰ and in female KO mice fed chow⁹⁰ or HFD,⁹¹ but in other studies, body fat was not increased in male KO mice fed chow^{90–93} or HFD⁹¹ and in female KO mice fed chow.^{91,92} Body fat was not increased in our male and female KO mice fed chow and in our male KO mice fed HFD. These data suggest that PYY deficiency plays little if any role in regulating mammalian body fat, consistent with the lack of obesity in humans with rare genetic PYY variants.⁵⁴

The GPCR FFAR4, a receptor for long chain free fatty acids (FFAs), purportedly mediates anti-obesity, anti-inflammatory and insulin sensitizing effects in vivo. In one study, BW was increased by 8 weeks of age in *Ffar4* KO mice fed HFD, consistent with increased obesity risk in humans with the inactivating *FFAR4* variant p.R270H.⁹⁴ However, BW was not increased in multiple studies of *Ffar* KO mice fed chow^{94,95} or HFD,^{95–97} consistent with lack of obesity in our HFD-fed *Ffar4* KO mice. Further, a FFAR4-selective agonist improved insulin sensitivity but not BW in WT mice and did not improve either parameter in *Ffar4* KO mice, and recent studies failed to link *FFAR4* variants, including p.R270H, to human obesity.^{98,99} These data suggest FFAR4 plays a minor, if any, role in regulating mammalian body fat.

Table 2 presents human gene expression and GWAS BMI data supporting the possibility that *GPR45*, *GPR61*, *KISS1*, *KISSIR*, *HDAC5*, *PRLHR*, *SLC6A4*, *SOST* and *OPRM1* variants contribute to obesity in humans as they do in mice.

Novel KO Lines

The obesity of KO mice in nine KO lines reported here, including *Kiss1* KO mice discussed above, is either a novel finding or was novel at the time the data were published by Lexicon Pharmaceuticals. KSR2 is a scaffold protein that likely coordinates signaling through kinase cascades. Our *Ksr2* KO and HET mice were obese,^{36,40,49} with 11/12 of HTS KO mice and 0/10 of HTS WT mice >2SD above the historic mean for %BF or BFg. These findings were confirmed in independent internal and external *Ksr2* KO models and in humans with *KSR2* haploinsufficiency;^{40,49,100} the human phenotype is consistent with human expression data and with a GWAS cluster of BMI variants that overlaps with the *KSR2* gene (Table 2). Here, we extend previous pair-feeding studies⁴⁰ to show that the obesity is due to both hyperphagia and decreased energy expenditure, and that KO mice survive long term only when pair-fed, with all KO mice dying soon after resuming ad-lib feeding (Supplementary Figure 2).

ASNSD1 is an understudied asparagine synthetase. *Asnsd1* KO mice have increased body fat and a progressive degenerative myopathy characterized by sarcopenia and myosteatosis;⁴⁵ HTS data showed 11/13 KO mice and 0/9 WT mice >2SD above the historic mean for %BF or BFg. *Asnsd1* KO mice have reduced viability (114 WT/251 HET/90 KO, $P<0.05$) that may be related to their

seizures and low systolic blood pressure. Additional studies are required to determine the contribution of myosteatosis to the increased body fat. The human *ASNSD1* expression pattern is consistent with a role for *ASNSD1* deficiency in human obesity and myopathy (Table 2).

G2E3 is a dual function ubiquitin ligase.¹⁰¹ We observed obesity and impaired glucose tolerance in multiple cohorts of *G2e3* KO mice;⁴⁷ HTS data showed 11/16 KO mice and 0/11 WT mice >2SD above the historic mean for %BF or BFg. These data are consistent with IMPC HTS data showing obesity in an independent *G2e3* KO model. The human gene resides 0.49 MBP 3' to a GWAS BMI variant cluster and is expressed in a pattern consistent with a role in body fat regulation (Table 2).^{102–104}

SRPK2 is a protein kinase overexpressed in many cancers.¹⁰⁵ We observed obesity in each of 7 small cohorts of *Srpk2* KO mice; when data were pooled and normalized, *Srpk2* KO mice showed a 41% increase in n%BF (Table 1), a 51% increase in nBFg (KO=151 ± 62 vs WT=100 ± 34, $P<0.001$ by Mann Whitney test) and 5% decrease in nLBM (KO=95 ± 13 vs WT=100 ± 9, $P<0.05$ by Mann Whitney test). In humans, only the *SRPK2* and *KMT2E* (lysine methyltransferase 2E) genes reside within a cluster of variants for waist–hip ratio adjusted BMI, a surrogate measure of abdominal adiposity;¹⁰⁶ these observations, and the *SRPK2* expression pattern, suggest that *SRPK2* may regulate visceral fat stores (Table 2). In cultured cells with mTORC1 activation, *SRPK2* knock-down inhibits de novo lipogenesis,¹⁰⁵ reinforcing the need to study additional *Srpk2* KO lines to confirm the phenotype reported here.

DPP8 is a cytoplasmic serine amino peptidase implicated in immune responses, cancer biology and cellular energy metabolism.^{107,108} Our *Dpp8* KO mice were obese, with a 33% increase in n%BF (Table 1), a 60% increase in nBFg (KO=160 ± 50 vs WT=100 ± 26, $P<0.001$), and a nonsignificant 5% increase in nLBM (KO=105 ± 15 vs WT=100 ± 7); HTS data showed 9/16 KO mice and 0/12 WT mice >2SD above the historic mean for %BF or BFg. *Dpp8* KO mice also had impaired glucose homeostasis; data pooled from 4 independent cohorts of *Dpp8* KO and WT mice showed that KO mice had fasting blood glucose and OGTT AUC values comparable to WT mice but had greater OGTT insulin levels at baseline and at 30 minutes (Supplementary Figure 3). In humans, *DPP8* is expressed in multiple tissues including adipose, hypothalamus, basal

ganglia and amygdala, and the human gene overlaps with a GWAS BMI cluster, consistent with a role in human body fat regulation (Table 2).

CXXC4, which can inhibit cancer cell growth by inhibiting WNT/β-catenin signaling, contains a CXXC-type zinc finger domain that can modify DNA methylation status.¹⁰⁹ Our *Cxxc4* KO mice were obese, with a 31% increase in n%BF (Table 1) and a 46% increase in nBFg (KO=146 ± 53 vs WT=100 ± 31, $P<0.001$ by Mann Whitney test). In humans, *CXXC4* is most highly expressed in hypothalamus (Table 2).

TENM3 and other teneurins are transmembrane proteins that participate in development of specific neuronal connectivity patterns.¹¹⁰ *Tenm3* KO mice had a 21% increase in n%BF (Table 1) and 24% increase in nBFg (KO=124 ± 43 vs WT=100 ± 34, $P<0.05$). An independent model of *Tenm3* KO mice had normal BW,¹¹¹ consistent with an insignificant 2% increase in BW of our *Tenm3* KO mice. In humans, *TENM3* is expressed in many tissues including adipose and in CNS, where it is highly expressed in hypothalamus (Table 2).

Obese KO Lines Studied with HTS Cohorts Only

KO Lines with Significantly Increased Body Fat That Support External Obese KO Lines

For many KO lines with significantly increased body fat in our HTS, confirmatory studies using additional cohorts were not performed, but our data support body fat data from external independent KO lines that were either published or evaluated in the IMPC HTS. Table 3 summarizes our data for these KO lines, and Table 4 presents human gene expression and/or GWAS data consistent with a role for the human orthologs of these targeted genes in regulating body fat.

ADCY3 and *NCOA1* genes are located near each other on human chromosome 2. Adenylyl cyclases are downstream enzymes for GPCRs, and *ADCY3* is a downstream effector for MC4R in hypothalamic neurons.¹¹² Our mice lacking one or both *Adcy3* alleles were obese (Table 3), which supported published findings in KO mice,^{113–115} in mice protected from DIO by gain-of-function *Adcy3* mutations,¹¹⁶ and in humans with *ADCY3* haploinsufficiency.^{117,118} The obesity observed in our *Ncoa1* KO mice is consistent with published KO data,¹¹⁹ with studies showing *NCOA1* modulates function of POMC neurons and energy homeostasis, and with human

Table 3 Body Fat Data for KO Lines with Significantly Increased Body Fat by HTS Only That Supports an External KO Line with High Body Fat

KO Line	Normalized % Body Fat						Normalized Body Fat (g)			
	Chow		HFD		Pooled Chow and HFD Data		Chow	HFD	Pooled Chow and HFD Data	
		KO/WT		KO/WT						
	Mean	(n/n)	Mean	(n/n)	KO	WT	Mean	Mean	KO	WT
<i>Adcy3</i> ^a	186	10/4	NM	NM	186 ± 84 ^{**†}	100 ± 12	240	NM	240 ± 184 ^{*†}	100 ± 22
<i>Aoc3</i>	120	8/4	NM	NM	120 ± 21	100 ± 5	133	NM	133 ± 27 ^{*†}	100 ± 4
<i>Apln</i>	121	8/4	125	6/4	122 ± 22 [*]	100 ± 20	124	142	131 ± 35 [*]	100 ± 36
<i>Ccn5</i>	129	8/4	120	6/6	125 ± 25 [*]	100 ± 31	125	108	118 ± 29	100 ± 43
<i>Ddah1</i>	106	8/4	134	8/8	134 ± 16 ^{*b}	100 ± 40	116	150	133 ± 45 ^b	100 ± 44
<i>Enox1</i>	116	8/4	126	6/5	120 ± 23 [*]	100 ± 15	127	128	127 ± 36	100 ± 18
<i>Herc1</i>	106	8/4	188	8/8	147 ± 49 ^{**†}	100 ± 23	106	235	169 ± 80 ^{*†}	100 ± 30
<i>Kdm3a</i>	150	8/4	NM	NM	150 ± 24 ^{**}	100 ± 25	162	NM	162 ± 42 [*]	100 ± 36
<i>Ncoa1</i>	110	8/4	120	8/8	115 ± 20	100 ± 22	119	130	125 ± 30 [*]	100 ± 32
<i>Ncs1</i>	119	8/4	131	6/6	124 ± 35 [*]	100 ± 28	128	148	137 ± 63	100 ± 38
<i>P2rx6</i>	131	8/4	NM	NM	131 ± 27 [*]	100 ± 7	162	NM	162 ± 51 ^{*†}	100 ± 4
<i>Pnpla2</i>	121	8/8	111	8/8	121 ± 20 ^{*†c}	100 ± 7	124	104	124 ± 30 ^{*†c}	100 ± 18
<i>Prmt7</i>	149	8/4	128	8/8	138 ± 32 ^{**}	100 ± 19	166	124	145 ± 57 ^{*†}	100 ± 27
<i>Retsat</i>	132	8/4	119	8/8	125 ± 32 [*]	100 ± 26	148	111	130 ± 51	100 ± 38
<i>Rgs10</i>	131	8/4	NM	NM	131 ± 19 ^{*†}	100 ± 4	150	NM	150 ± 43 ^{*†}	100 ± 15
<i>Tnfrsf13b</i>	126	8/4	NM	NM	126 ± 28	100 ± 13	141	159	141 ± 36 ^{*†}	100 ± 16
<i>Usp38</i>	142	8/4	133	8/7	138 ± 33 ^{**}	100 ± 21	159	140	149 ± 52 ^{**}	100 ± 28

Notes: KO mice different from WT mice, **P* < 0.05; ***P* < 0.01; † Statistical analysis by Mann–Whitney test. ^aKO data represent data pooled from 2 KO and 8 HET *Adcy3* mice. ^bHFD data only. ^cChow data only.

Abbreviations: HFD, high fat diet; (g), grams; KO, knockout; WT, wild-type; n, number of mice; NM, not measured.

data linking *NCOA1* inactivation to obesity.¹²⁰ These human data suggest that variants linked to both *ADCY3* and *NCOA1* contribute to the strong and broad GWAS BMI cluster that overlaps these genes (Table 4).

Mice with KOs of genes encoding the secreted proteins TNFSF13B,^{121,122} CCN5,¹²³ and APLN¹²⁴ are obese, consistent with data from our independent KO lines (Table 3). Of note, a third *Ccn5* KO line was not obese in the IMPC HTS, which may be consistent with the modest obesity observed in the published,¹²³ and our (Table 3), KO lines.

Two independent KOs of *Kdm3a*, encoding a histone demethylase, were obese,^{125–127} findings supported by HTS data from an additional IMPC KO line and from our KO mice (Table 3), where 4/8 KO mice and 0/4 WT mice were >2SD above the historic mean for %BF or BFg. GWAS data show *KDM3A* sharing a BMI cluster with 2 genes, *CHMP3* (charged multivesicular body protein 3) and *RNF103* (ring finger protein 103), that lack an obvious link to obesity.

The IMPC HTS reported that KOs of *Herc1* and *Usp38*, which encode enzymes involved in ubiquitination pathways, were obese, consistent with our data (Table 3); body composition was not evaluated in two *Usp38* KO

publications.^{128,129} Although some human *HERC1* variants are associated with somatic overgrowth, they result in increased, not decreased, *HERC1* activation and thus are not relevant here.^{130,131} Both *HERC1* and *USP38* reside near BMI clusters in gene-poor chromosomal regions (Table 4), suggesting a possible link to human obesity for each gene.

Mice with a KO of *Prmt7*, which influences skeletal muscle oxidative metabolism, were obese;¹³² HTS data on our KO line (Table 3) and an independent KO line from the IMPC support this finding. GWAS data show *PRMT7* as the only obesity-linked gene among 5 genes located within in a BMI cluster (Table 4). The published finding that *Retsat* KO mice were obese¹³³ is confirmed by our KO data (Table 3).

Inactivating mutations in the lipase *PNPLA2* result in neutral lipid storage disease with myopathy (NLSDM) in humans and in mice,^{134–137} with modest obesity observed in chow-fed, but not HFD-fed, *Pnpla2* KO mice. Our *Pnpla2* KO mice also had NLSDM (data not shown), and the increased body fat data presented in Table 3 are from chow-fed mice only because our KO mice were not

Table 4 Human Expression and GWAS Data for KO Lines with Significantly Increased Body Fat by HTS Only That Supports an External KO Line with High Body Fat

KO Line	Human Expression ^a	Major GWAS BMI Clusters Within 1.2 MBP ^b		
		Lead Variant	P value	Cluster Distance
<i>Adcy3</i>	Wide; CNS, incl Hyp; Ad	2:25141538:A:G 2:25378372:G:A*	<e-109 <e-20	Overlapping Overlapping
<i>Aoc3</i>	High Ad; no CNS	17:41058634:C:T* 17:41353410:A:G	<e-10 <e-9	Overlapping 0.08 MBP 3'
<i>Apln</i>	Wide; CNS, incl Hyp; low Ad	X:128143072:G:A**	<e-9	0.64 MBP 3'
<i>Ccn5</i>	Mainly Ad	None		
<i>Ddah1</i>	Wide; CNS, incl Hyp; Ad	1:86264061:G:C*	<e-13	<0.01 MBP 5'
<i>Enox1</i>	Wide; CNS, incl Hyp; Ad	None		
<i>Herc1</i>	Wide; CNS, incl Hyp; Ad	15:63868761:A:G* 15:63789952:A:C	<e-9 <e-10	Overlapping 0.1 MBP 3'
<i>Kdm3a</i>	Wide; CNS, incl Hyp; Ad	2:86764004:C:T	<e-13	Overlapping
<i>Ncoa1</i>	Wide; CNS, incl Hyp; Ad	2:25141538:A:G 2:25378372:G:A*	<e-109 <e-20	Overlapping Overlapping
<i>Ncs1</i>	Wide; CNS, incl Hyp; low Ad	9:133787225:A:G	<e-13	0.78 MBP 3'
<i>P2rx6</i>	Wide; CNS, incl Hyp; Ad	22:22190163:C:A	<5e-8	0.81 MBP 3'
<i>Pnpla2</i>	Mainly Ad	11:817786:C:T* 11:891338:G:A	<e-15 <e-11	Overlapping Overlapping
<i>Prmt7</i>	Wide; CNS, incl Hyp; Ad	16:68381978:A:G 16:67409180:G:A*	<e-15 <e-14	Overlapping 0.94 MBP 5'
<i>Retsat</i>	High Ad; low CNS, incl Hyp	2:86764004:C:T	<e-13	1 MBP 5'
<i>Rgs10</i>	Wide; low CNS, incl Hyp; Ad	10:120465796:C:G	<e-10	0.75 MBP 3'
<i>Tnfrsf13b</i>	Whole blood; Spleen; Lung	13:107882445:A:G	<e-10	1.02 MBP 5'
<i>Usp38</i>	Wide; CNS, incl Hyp; Ad	4:143663206:T:C	<5e-8	0.37 MBP 5'

Notes: ^aData obtained from <https://gtexportal.org/home/>. ^bData obtained from <https://hugeamp.org/> on 8/17/2021. *Waist-hip ratio adjusted BMI; **Weight GWAS variants only.
Abbreviations: MBP, million base pairs; incl, including; CNS, central nervous syndrome; Hyp, hypothalamus; Ad, adipose.

obese when fed HFD. *Ddah1* KO mice had increased BW on HFD but not on chow,¹³⁸ consistent with our findings (Table 3); consistent with these findings, chow-fed *Ddah1* KO mice were not obese in the IMPC HTS.

Mice with a KO of *Aoc3* which encodes VAP-1/SSAO are modestly obese,^{139,140} consistent with our findings (Table 3); interestingly, the human *AOC3* gene is expressed in adipose tissue but not CNS and resides within a waist-hip ratio adjusted BMI cluster (Table 4), suggesting a role for AOC3 in regulating visceral fat mass. *Enox1* is a little-studied gene,¹⁴¹ and the finding that *Enox1* HET mice were obese in the IMPC HTS is consistent with the obesity observed in our *Enox1* KO mice (Table 3).

External KOs for the neuronal calcium sensor *Ncs1* and the GTPase-activating protein *Rgs10* were obese,^{142,143} consistent with our findings (Table 3). Although no publications link the P2RX6 channel¹⁴⁴ to obesity, the IMPC HTS found that male *P2rx6* KO mice were obese; the obesity observed in our *P2rx6* KO mice (Table 3) was present in both males and females (data not shown).

KO Lines with Numerically Increased Body Fat That Support External Obese KO Lines

For many KO lines with a numerical, but not statistically significant, body fat increase of at least 15% in our HTS, confirmatory studies using additional cohorts were not

Table 5 Body Fat Data for KO Lines with Numerically Increased Body Fat by HTS Only That Supports an External KO Line with High Body Fat

KO Line	Normalized % Body Fat						Normalized Body Fat (g)			
	Chow		HFD		Pooled Chow and HFD Data		Chow	HFD	Pooled Chow and HFD Data	
		KO/WT		KO/WT						
	Mean	(n/n)	Mean	(n/n)	KO	WT	Mean	Mean	KO	WT
<i>Adamts18</i>	113	8/4	126	6/6	119 ± 27	100 ± 31	103	121	111 ± 36	100 ± 43
<i>Adm2</i>	138	8/4	111	4/4	129 ± 53	100 ± 14	181	126	163 ± 115	100 ± 14
<i>Aqp7</i>	117	8/4	NM	NM	117 ± 26	100 ± 8	127	NM	127 ± 36	100 ± 10
<i>AU040320</i>	113	8/4	119	8/8	116 ± 22	100 ± 26	126	128	127 ± 39	100 ± 36
<i>Glr2</i>	110	8/4	127	8/8	119 ± 32	100 ± 26	109	137	123 ± 44	100 ± 30
<i>Gpx7</i>	124	8/4	123	7/7	124 ± 37	100 ± 21	138	128	134 ± 61	100 ± 24
<i>Hdac6</i>	116	8/4	112	6/6	114 ± 26	100 ± 12	153	117	137 ± 50	100 ± 20
<i>Igf2</i>	115	8/4	121	5/5	117 ± 19	100 ± 25	122	126	124 ± 31	100 ± 28
<i>Npyf</i>	105	8/4	139	6/5	119 ± 31	100 ± 21	101	155	125 ± 53	100 ± 41
<i>Nr4a1</i>	116	8/4	132	8/8	124 ± 31	100 ± 40	122	142	132 ± 40	100 ± 54
<i>Prdx6</i>	105	8/4	119	6/6	111 ± 23	100 ± 9	123	124	124 ± 42	100 ± 15
<i>St3gal2</i>	106	8/4	138	5/5	123 ± 31	100 ± 18	113	136	119 ± 35	100 ± 21
<i>Usp13</i>	108	8/4	125	4/4	114 ± 18	100 ± 12	112	132	119 ± 25	100 ± 17

Abbreviations: HFD, high fat diet; (g), grams; KO, knockout; WT, wild-type; n, number of mice; NM, not measured.

performed but our data support external obesity data. Table 5 summarizes our data for these KO lines, and Table 6 presents human gene expression and/or BMI GWAS data consistent with a role for the human orthologs of these targeted genes in regulating body fat.

There are 4 secreted proteins in this category. Our HTS data (Table 5) were consistent with the modest obesity observed in *Igf2* KO mice at 8 weeks of age¹⁴⁵ and with IMPC HTS data showing increased body fat in *AU040320* KO mice. Male infertility, but no body fat data, was reported in an independent *AU040320* KO line,¹⁴⁶ consistent with the male infertility of KO mice studied by both the IMPC and Lexicon. Our HTS data were also consistent with the ability of ADM2 to inhibit HFD-induced obesity when overexpressed by adipose tissue of transgenic mice^{147,148} and to inhibit food intake when delivered subcutaneously to WT mice,¹⁴⁹ and with the increased BW of *Adm2* HET mice.¹⁵⁰ Central delivery of NPVF, encoded by a gene expressed only in hypothalamus (Table 6), induced acute anorexia in chicks,^{151,152} consistent with our *Npyf* KO data; of interest, body fat was significantly increased only in our KO mice fed HFD (nBF: KO 157 ± 52 vs WT 100 ± 35, n%BF: KO 139 ± 21 vs WT 100 ± 25, both $P < 0.05$), suggesting that NPVF may regulate intake of palatable diets.

For the 7 enzymes in this category, obesity of KO mice from published *St3gal2*, *Glr2*, *Hdac6* and *Prdx6* KO lines

agrees with our data.^{153–157} Obesity of KO mice from published *Gpx7* and *Adamts18* KO lines^{158,159} is consistent with our data and with GWAS data linking obesity to *GPX7*¹⁵⁸ and to *ADAMTS18* which is closely associated with a BMI cluster (Table 6), but not with the absence of obesity in independent *Gpx7* and *Adamts18* KO lines studied in the IMPC HTS. Our HTS data for *Usp13* KO mice are consistent with the obesity of female mice from an independent IMPC *Usp13* KO line.

The obesity of mice with a KO of the nuclear hormone receptor *Nr4a1*^{160,161} is consistent with our HTS data. AQP7 is the primary glycerol transporter for adipocytes and cardiomyocytes. Our HTS data support the obesity observed in published cohorts of *Aqp7* KO mice.^{162,163} We also observed focal myocardial degeneration in 3 of 4 KO mice (Supplementary Figure 4), consistent with published data showing impaired myocardial response to pressure overload in *Aqp7* KO mice.¹⁶⁴

KO Lines with Significantly Increased Body Fat and No External Obese KO Lines

For many KO lines with significantly increased body fat in our HTS, confirmatory studies examining additional cohorts were not performed and no supportive external KO data are available. We have many KO lines in this category, but our presentation here is focused on data from KO lines (Table 7) where the human ortholog of the

Table 6 Human Expression and GWAS Data for KO Lines with Numerically Increased Body Fat by HTS Only That Supports an External KO Line with High Body Fat

KO Line	Human Expression ^a	Major GWAS BMI Clusters Within 1.2 MBP ^b		
		Lead Variant	P value	Cluster Distance
<i>Adamts18</i>	Mainly Ad and CNS, incl Hyp	16:77255496:A:C	<e-9	0.03 MBP 3'
<i>Adm2</i>	Thyroid, Kidney, GI; no CNS, Ad	22:50709495:A:G	<e-9	0.2 MBP 5'
<i>Aqp7</i>	Mainly Ad; Heart; not in Hyp	9:33921919:T:C	<e-11	0.23 MBP 5'
<i>AU040320</i>	Wide; CNS, incl Hyp; Ad	None		
<i>Glrx2</i>	Wide; CNS, incl Hyp; Ad	1:193024644:C:G	<5e-8	0.02 MBP 3'
<i>Gpx7</i>	Wide; high Ad; CNS low, incl Hyp	None		
<i>Hdac6</i>	Wide; CNS, incl Hyp; Ad	None		
<i>Igfbp2</i>	High vAd; Liver; not in CNS	2:218393389:G:A*	<e-11	0.82 MBP 3'
<i>Npvf</i>	Hyp only	7:25871109:C:T*	<e-71	0.56 MBP 5'
		7:24325009:G:A	<e-14	0.78 MBP 3'
<i>Nr4a1</i>	Wide; high Ad; CNS low, incl Hyp	None		
<i>Prdx6</i>	Wide; Ad; low CNS, incl Hyp	1:174063646:A:G	<e-16	Overlapping
		1:172352990:G:A*	<e-54	0.99 MBP 5'
<i>St3gal2</i>	Wide; CNS, incl Hyp; Ad	16:69556715:C:T	<e-26	Overlapping
<i>Usp13</i>	SkM; Ad; low CNS, incl Hyp	3:180736253:T:C	<e-11	0.87 MBP 3'

Notes: ^aData obtained from <https://gtexportal.org/home/>. ^bData obtained from <https://hugeamp.org/> on 8/17/2021. *Waist-hip ratio adjusted BMI.

Abbreviations: MBP, million base pairs; incl, including; CNS, central nervous syndrome; Hyp, hypothalamus; Ad, adipose; GI, gastrointestinal tract; vAd, visceral adipose; SkM, skeletal muscle.

targeted gene is closely associated with a GWAS BMI variant cluster and has an expression pattern consistent with a role for the ortholog in regulating body fat.

Tle4 KO mice showed reduced viability and poor growth in the first few weeks of life.^{165,166} At weaning, our *Tle4* mice had a Mendelian ratio of 37 WT, 78 HET and 24 KO mice that trends toward reduced viability ($P=0.1$), and BWs of male and female KO mice were lower than those of WT littermates. However, by 8 weeks of age, BWs of KO mice had caught up to or surpassed those of WT littermates (Supplementary Figure 5). This trend continued as the mice aged. Unlike body composition studies of our other HTS KO lines, *Tle4* KO mice were older with ages ranging from 19 to 52 weeks; nevertheless, the %BF and BFG for all 6 *Tle4* WT mice were within 1 SD of the HTS historic mean, while 13/14 KO mice were >2SD above the historic mean for %BF and BFG. Blood glucose and total cholesterol levels were also higher for KO vs WT littermate mice: for blood glucose, KO = 142 ± 35 (n=19) vs WT = 97 ± 27 mg/dL (n=7), $P<0.01$; for total

cholesterol, KO = 146 ± 51 (n=11) vs WT = 72 ± 19 mg/dL (n=2), $P<0.05$ by Mann-Whitney test. In humans, GWAS BMI clusters are found within and 0.81 MBP 5' to the *TLE4* gene, which resides alone in the middle of a 2.5 MBP stretch of chromosome 9 (Table 8), and a T2D cluster (lead variant 9:81908842:T:C, $P<e-22$) is found 0.23 MBP 5' to the *TLE4* gene (<https://hugeamp.org/>), further supporting a role for *TLE4* in obesity and T2D.

No external KOs or published links to obesity were identified for the kinase *AK5*,¹⁶⁷ but a strong GWAS BMI cluster is found within, and is primarily associated with, the human *AK5* gene that is expressed almost exclusively in the CNS (Table 8). No body composition data were presented for an external KO of the cell adhesion molecule encoded by *Ntm*,¹⁶⁸ but two independent GWAS BMI clusters are found within, and are only associated with, the human *NTM* gene (Table 8). Perinatal lethality occurred in the IMPC KO for *Tusc3*, which encodes a subunit of the oligosaccharyl transferase responsible for N-glycosylation of nascent proteins;¹⁶⁹ no published links to obesity were found for *Tusc3* but a BMI cluster is

Table 7 Body Fat Data for KO Lines with Significantly Increased Body Fat by HTS and No External KO Line with High Body Fat

KO Line	Normalized % Body Fat						Normalized Body Fat (g)			
	Chow		HFD		Pooled Chow and HFD Data		Chow	HFD	Pooled Chow and HFD Data	
		KO/WT		KO/WT						
	Mean	(n/n)	Mean	(n/n)	KO	WT	Mean	Mean	KO	WT
<i>Adams4</i> ^a	119	36/12	NM	NM	119 ± 27**†	100 ± 13	137	NM	137 ± 44***†	100 ± 17
<i>Ak5</i>	136	8/4	135	8/8	135 ± 29**	100 ± 15	183	166	173 ± 48***†	100 ± 15
<i>Ankk1</i>	126	8/4	135	8/8	129 ± 34*	100 ± 33	138	148	144 ± 59*	100 ± 40
<i>Dgkg</i>	118	8/4	123	7/7	121 ± 31	100 ± 19	127	139	133 ± 45*	100 ± 29
<i>Igdcc4</i>	120	8/4	177	8/8	149 ± 49*	100 ± 41	126	176	150 ± 59*	100 ± 57
<i>Itih1</i>	120	8/4	110	8/8	115 ± 18*	100 ± 20	120	129	125 ± 32*	100 ± 29
<i>Lrrn2</i>	157	8/4	123	8/8	141 ± 46**	100 ± 14	175	125	151 ± 47*	100 ± 18
<i>Lrrtm1</i>	164	8/4	105	4/4	144 ± 42*	100 ± 23	170	104	150 ± 52*	100 ± 31
<i>Mfap3l</i>	135	8/4	136	7/7	135 ± 26**	100 ± 29	140	148	144 ± 42**	100 ± 31
<i>Ntm</i>	128	8/4	131	7/7	129 ± 17***	100 ± 17	125	132	128 ± 28*	100 ± 27
<i>Pecr</i>	139	8/4	131	6/3	136 ± 42*†	100 ± 16	152	120	138 ± 62*†	100 ± 17
<i>Prok2</i> ^b	154	5/4	NM	NM	154 ± 100	100 ± 12	129	NM	129 ± 108	100 ± 18
<i>Prokr2</i>	161	8/4	NM	NM	161 ± 23***	100 ± 7	121	NM	121 ± 27	100 ± 10
<i>Ptp4a1</i>	126	8/4	137	7/9	131 ± 19***	100 ± 15	129	152	140 ± 33**	100 ± 18
<i>Ptprn</i>	121	8/4	NM	NM	121 ± 27	100 ± 10	144	NM	144 ± 34*	100 ± 11
<i>Resp18</i>	107	8/4	131	8/8	119 ± 24*	100 ± 23	110	159	134 ± 39*	100 ± 35
<i>Scg3</i> ^c	131	28/11	101	6/6	126 ± 36**†	100 ± 22	139	100	132 ± 52*†	100 ± 21
<i>Sik2</i>	135	8/4	140	7/7	137 ± 31**	100 ± 27	146	157	152 ± 57*	100 ± 36
<i>Tle4</i> ^d	189	14/6	NM	NM	189 ± 41***†	100 ± 14	312	NM	312 ± 130***†	100 ± 25
<i>Tusc3</i>	134	8/4	115	6/6	126 ± 29	100 ± 35	149	122	137 ± 42*	100 ± 42
<i>Wnt8b</i>	149	8/4	137	8/8	144 ± 37**	100 ± 27	172	143	157 ± 54**	100 ± 39

Notes: KO mice different from WT mice, **P* < 0.05; ***P* < 0.01; ****P* < 0.001; † Statistical analysis by Mann–Whitney test. ^aData pooled from 3 independent *Adams4* KO lines, 2 targeting exon 4 and 1 gene trap in the intron between coding exons 1 and 2; pooled data from 24 KO and 12 HET mice compared to pooled data from 12 WT mice. ^bDifferences not statistically significant, but *Prok2* data included due to similarity with receptor *Prokr2*. ^cData pooled from 2 independent *Scg3* KO lines, 1 targeting exons 1–4 and 1 gene trap in the intron between the last 2 coding exons; pooled data from 22 KO and 12 HET mice compared to pooled data from 17 WT mice. ^dAge of *Tle4* KO and WT littermate mice ranged from 19 to 52 weeks.

Abbreviations: HFD, high fat diet; (g), grams; KO, knockout; WT, wild-type; n, number of mice; NM, not measured.

located within, and only associated with, the human *TUSC3* gene (Table 8).

There are no external *Ankk1* KO lines. The human Taq1A RFLP (rs1800497) linked to the dopamine 2 receptor (DRD2) causes a single amino acid substitution in the ANKK1 substrate-binding domain.¹⁷⁰ The A1 allele is associated with 1) obesity,¹⁷¹ 2) decreased striatal DRD2 density¹⁷² and 3) increased food intake.¹⁷³ This suggests that, if the A1 allele expresses a less active ANNK1 form, *Ankk1* KO mice will be obese,¹⁷⁴ a hypothesis consistent with our findings.

A few of our KO lines target genes having human orthologs that share GWAS BMI clusters with other genes that have no published or IMPC HTS data linking them to obesity. No external KO was identified for the cell adhesion protein *Igdcc4*, which is linked to early childhood adiposity,¹⁷⁵ *IGDCC4* is located in a BMI cluster

(Table 8) shared with *DPP8*, discussed above, and with 5 other genes with no clear link to obesity. No external KO was identified for *Itih1*, which encodes a heavy chain for the serine protease inhibitor inter-alpha trypsin inhibitor. Although ITIH1 is expressed only in liver, studies link this secreted protein to mood disorders and report that overexpressed ITIH1 can deposit on hyaluronan surrounding mouse adipose tissue.^{176,177} *ITIH1* is centrally located in, and the landmark for,¹⁰⁴ a large BMI cluster which overlaps 14 additional genes, none of which have been linked to obesity. Although external *Lrrtm1* KO lines had normal BW,^{178,179} and body fat was not increased in the IMPC HTS, this KO line is included because of the robust body fat phenotype with 5/12 KO mice and 0/8 WT mice >2SD above the historic mean for %BF or BFG, and because *LRRTM1* and *CTNNA2* (catenin alpha 2) are the only genes sharing a BMI cluster (Table 8). No external KOs

Table 8 Human Expression and GWAS Data for KO Lines with Significantly Increased Body Fat by HTS Only and No External KO Line with High Body

KO Line	Human Expression ^a	Major GWAS BMI Clusters Within 1.2 MBP ^b		
		Lead Variant	P value	Cluster Distance
<i>Adamts4</i>	vAd highest; low CNS, incl Hyp	1:160412880:G:A*	<e-11	0.74 MBP 3'
<i>Ak5</i>	Mainly CNS, incl Hyp	1:77967507:T:A	<e-35	Overlapping
<i>Ankk1</i>	Wide; CNS, incl Hyp; Ad	11:113271360:C:T	<5e-8	Overlapping
<i>Dgkg</i>	Wide; CNS, incl Hyp; Ad	3:185834499:A:T	<e-66	Overlapping
<i>Igdcc4</i>	Wide; CNS, incl Hyp; Ad	15:65916527:A:T	<5e-8	0.08 MBP 5'
<i>Itih1</i>	Liver-specific	3:5255316:G:A* 3:52740182:C:G	<e-45 <e-41	Overlapping Overlapping
<i>Lrrn2</i>	Wide; CNS, Hyp highest; high vAd	1:204414769:T:C* 1:203491392:T:G	<5e-8 <e-11	0.17 MBP 3' 1.1 MBP 3'
<i>Lrrtm1</i>	Mainly CNS, incl Hyp	2:80456138:T:C	<5e-8	Overlapping
<i>Mfap3l</i>	Wide; highest in CNS, incl Hyp; Ad	4:170955277:G:T	<e-9	Overlapping
<i>Ntm</i>	Mainly CNS, incl Hyp; low Ad	11:131452912:T:C 11:131957293:T:C 11:130877142:T:C	<e-11 <e-10 <e-15	Within NTM Within NTM 0.35 MBP 5'
<i>Pecr</i>	Wide; high Ad; lower CNS, incl Hyp	2:216904019:A:T 2:216295312:T:A*	<5e-8 <e-13	Within PECCR 0.55 MBP 3'
<i>Prok2</i>	Mainly whole blood; Lung; Spleen	3:71668037:T:C	<5e-8	0.15 MBP 3'
<i>Prokr2</i>	Mainly CNS, incl Hyp; Pit; no Ad	20:5667459:T:C*	<e-28	0.35 MBP 5'
<i>Ptp4a1</i>	Wide; CNS, incl Hyp; Ad	6:64247773:G:C	<e-10	Overlapping
<i>Ptprn</i>	Wide; CNS, Hyp highest; Pit	2:219903258:T:G	<e-14	Overlapping
<i>Resp18</i>	CNS-specific, incl Hyp	2:219903258:T:G	<e-14	Overlapping
<i>Scg3</i>	Mainly CNS, incl Hyp; Pit; no Ad	15:51754451:T:C	<e-12	Overlapping
<i>Sik2</i>	Wide, incl CNS; highest in Ad	11:111642119:G:A*	<e-34	Overlapping
<i>Tle4</i>	Wide; CNS, incl Hyp; Ad	9:82229981:T:G 9:81341229:G:T	<e-10 <e-11	Within TLE4 0.81 MBP 5'
<i>Tusc3</i>	Wide; CNS, Hyp highest; Ad	8:15565257:T:C	<e-10	Within Tusc3
<i>Wnt8b</i>	Wide; CNS, incl Hyp; Ad	10:102395440:T:C	<e-20	0.15 MBP 3'

Notes: ^aData obtained from <https://gtexportal.org/home/>. ^bData obtained from <https://hugeamp.org/> on 8/17/2021. *Waist-hip ratio adjusted BMI.

Abbreviations: MBP, million base pairs; incl, including; CNS, central nervous syndrome; Hyp, hypothalamus; Ad, adipose; vAd, visceral adipose; Pit, pituitary.

or published links to obesity were identified for the gene encoding the protein kinase *Mfap3l*; ¹⁸⁰ *MFAP3L* and *AADAT* (aminoadipate aminotransferase) are the only genes sharing a BMI cluster (Table 8). No external KOs or published links to obesity were identified for the gene encoding the short-chain dehydrogenase/reductase *PECR*; ¹⁸¹ *PECR* and *MREG* (melanoregulin) share a BMI cluster and are the two closest genes to a body fat

percentage cluster (lead variant 2:216661156:G:C, $P < e-21$) located 0.2 MBP 3' to the *PECCR* gene (<https://hugeamp.org/>). Although one external KO line for the gene encoding the protein tyrosine phosphatase *Ptp4a1* ¹⁸² did not evaluate body fat and a second showed normal body fat in the IMPC HTS, support for our finding of obese KO mice was provided by the link between *PTP4A1* and addictive behaviors ¹⁸³ and by a BMI cluster

that overlaps *PTP4A1* and is shared only by *PHF3* (PHD finger protein 3) and *EYS* (eyes shut homolog) (Table 8). *SCG3* is a protein that is co-expressed with, and forms secretory granules with, several appetite-regulating peptides in the hypothalamus; analysis of human genetic variants suggested that decreased *SCG3* expression increases obesity risk.^{8,184} Although an independent IMPC line of *Scg3* KO mice was not obese, our finding of obese KO and HET mice in 2 independent *Scg3* KO lines (Table 7), one targeted by homologous recombination and the other by gene trapping (Supplementary Tables 2 and 3), and the observation that *SCG3* resides within a BMI cluster (Table 8) that is shared with *DMXL2* (Dmx like 2) which has no clear link to obesity, support a role for *SCG3* in the regulation of body fat.

RESP18 and *PTPRN* are intracellular proteins that participate in hormone secretion pathways; the genes for these proteins, which share ~40% amino acid similarity over a 200 amino acid stretch of *PTPRN*, are located adjacent to each other in a head-to-tail orientation in mammalian genomes, suggesting they are evolutionarily related.^{185,186} *Resp18*, first identified as a gene coregulated with *POMC*, is downregulated by dopamine and is highly expressed in the diencephalon, suggesting a role in salt and water balance and/or feeding behavior,¹⁸⁵ indeed, a rat *Resp18* KO line, although not evaluated for body fat, did show increased SBP.¹⁸⁷ *PTPRN*, also known as IA-2, is an autoantigen linked to T1D but not obesity, and published *Ptpn* KO mice have normal BW.¹⁸⁶ Surprisingly, KO lines for both genes had increased body fat in our agnostic screen, and in addition *Resp18* KO mice had increased SBP (KO, $n=8$, 101 ± 7 vs WT, $n=4$, 92 ± 7 mmHg, $P<0.05$) supporting the finding in *Resp18* KO rats. *PTPRN* and *RESP18* are the only 2 of 8 genes located in a BMI cluster (Table 8) that are currently linked to obesity.

A few of our KO lines target genes having human orthologs that share a GWAS BMI cluster with other genes linked to obesity. Genes for the kinase *Dgk*¹⁸⁸ and the neuronal membrane protein *Lrrn2*¹⁸⁹ have no publicly available KO data or published links to obesity despite having increased body fat in our HTS (Table 7). *DGKG* and *ETV5* (ETS variant transcription factor 5) are the only genes sharing a BMI cluster on human chromosome 3, while *LRRN2* and *MDM4* (MDM4 regulator of p53) are the only genes sharing a BMI cluster on human chromosome 1 (Table 8); the low body fat observed in *Etv5* KO mice¹⁹⁰ and *Mdmx* KO mice¹⁹¹ suggests that one or both genes in each of these two BMI clusters may

contribute to the GWAS signal. A published KO of *Wnt8b*¹⁹² did not provide body fat data and the limited BW data did not support the finding of obesity that we observed in our KO line (Table 7). *WNT8B* shares a nearby BMI cluster (Table 8) with 4 other genes including *HIF1AN* (hypoxia inducible factor 1 subunit alpha inhibitor); the low body fat observed in *Hif1an* KO mice¹⁹³ suggests that multiple genes might contribute to this GWAS cluster. External *Sik2* KO mice had impaired glucose homeostasis and increased circulating TG and adipocyte size consistent with a role for *SIK2* in adipogenesis, but body fat was not increased.¹⁹⁴ Our *Sik2* KO mice had an insignificant 23% increase in OGTT glucose AUC and a significant 9% increase in HbA1c with KO = $4.8\pm0.4\%$ ($n=8$) vs WT = $4.4\pm0.2\%$ ($n=4$), $P<0.05$, consistent with modestly impaired glucose tolerance observed in the published *Sik2* KO mice. However, our KO mice also had increased body fat, consistent with high expression of the human *SIK2* gene in adipose tissue and the association of this gene with an overlapping waist-hip ratio adjusted BMI cluster (Table 8). This BMI cluster is shared with 11 genes including *CRYAB*; the possible link of *CRYAB* to adiposity¹⁹⁵ suggests that multiple genes might contribute to this GWAS BMI signal.

In humans, *PROK2* is a CNS peptide that binds the GPCR *PROKR2*; inactivating mutations of either gene can result in Kallmann Syndrome, the combination of hypogonadotropic hypogonadism and anosmia.¹⁹⁶ Consistent with these findings, *Prok2* and *Prokr2* KO mice have hypoplastic olfactory bulbs and atrophic reproductive organs.^{197,198} Our *Prok2* and *Prokr2* KO mice confirm these findings, including male and female infertility for *Prok2* KO mice, but likely also share an obesity phenotype not previously reported; although our KO mouse numbers were small, 7/8 *Prok2* and 3/5 *Prokr2* KO mice were >2SD above the historic mean for %BF, compared to 0 WT mice for either cohort. These data are consistent with the ability of *PROK2*, a hypothalamic peptide, to inhibit food intake and promote weight loss when administered to rats.¹⁹⁹ Recent data suggest this link to obesity may also extend to humans.²⁰⁰

Published and IMPC HTS data describing external KOs for the secreted protein *Adamts4*²⁰¹ did not provide body fat or BW data supporting the obesity of our mice (Table 7), and the human ortholog is not closely associated with a BMI cluster (Table 8). *Adamts4* was included because KO and HET mice from each of our 3 independent KO lines, two targeting exon 4 by homologous

recombination and one the result of a gene trapping vector introduced into the intron between coding exons 1 and 2 (Supplementary Tables 2 and 3), showed similar increases in both body fat (pooled data in Table 7) and in total cholesterol, where data pooled from all KO (n=24) and HET (n=12) mice = 143 ± 31 vs data pooled from all WT mice (n=12) = 118 ± 30 mg/dL, $P < 0.05$.

Discussion

We provide data for multiple mouse KO models with established and novel high body fat phenotypes. To better understand the potential value of these observations, we considered 1) if there is fidelity between mouse and human obesity genes; 2) if there is value in identifying obesity genes; and 3) criteria for what constitutes confirmation of a novel mouse KO obesity phenotype.

The High Fidelity Between Mouse and Human Obesity Genes

Mice are considered a good model for human disease,^{26,202} consistent with our finding that skeletal and other phenotypes are highly conserved between mice and man.^{28,37} Nevertheless, the limited ability of mouse models to mimic human inflammatory diseases²⁰³ demands a thorough analysis of the fidelity between mouse and human obesity genes. As shown in Table 9, inactivating mutations in many human genes are sufficient by themselves to result in obesity; for our purposes here, these are referred to as monogenic obesity genes.^{40,117,118,120,204–233} For each of these 18 human monogenic obesity genes, inactivating mutations in the orthologous mouse gene closely reproduce the human phenotype;^{10–13,36,49,59–62,100,114,115,119,220,232,234–261} in fact, 13 of these genes were initially reported as monogenic obesity genes in mice, including *Lep*,^{10,204} *Lepr*,^{10,207} *Mc4r*,^{13,209} *Pomc*,^{212,235} *Sh2b1*,^{214,238} *Bdnf*,^{216,241} *Ntrk2*,^{217,243} *Ksr2*,^{36,40} *Adcy3*,^{114,117,118} *Cpe*,^{222,249} *Tub*,^{223,249} *Ncoa1*^{119,120} and *Mc3r*.^{60,61,227,256} In addition, *Mrap2*,²²⁰ *Cep19*²²⁴ and *Cartpt*^{225,253} were simultaneously reported as obesity genes in mice and humans. Only *Sim1*^{218,245} and *Pcskl*^{228,232,257–261} were first reported as obesity genes in humans. For *Mc3r*, KO mice are clearly obese with decreased LBM^{36,60–62} but evidence in humans was weak until recent studies focused on the human C17A+G241A haplotype encoding a partially inactivated receptor that, when homozygous, is associated with increased fat mass and decreased LBM.^{226,227} This phenotype was recently

reproduced in mice homozygous for the human *MC3R* containing this haplotype compared to mice homozygous for WT human *MC3R*; of interest, this mouse phenotype is less severe than that observed in *Mc3r* KO mice.^{226,256}

PCSK1 encodes proprotein convertase 1/3 (PC1/3), an enzyme that processes precursor neuropeptides and pro-hormones in endocrine tissues. Humans with complete deficiency develop severe obesity and a complex set of endocrinopathies,^{228,229} while *Pcskl* KO mice that survive to adulthood are not obese,²⁵⁷ which suggests that PC1/3 enzymatic pathways regulating body fat in humans are not operative in mice. However, humans heterozygous for the null mutation or with mutations causing partial loss of PC1/3 function develop obesity,^{230–233} a finding duplicated in HET mice and in mice with partial loss of PC1/3 activity,^{257–261} suggesting that the PC1/3 enzymatic pathways regulating body fat in humans are indeed shared by mice. Thus, the available evidence indicates a remarkable conservation between mice and man of the genes that regulate body fat.

Only 5 of the 72 obese mouse KO lines reported here appear on this list of shared mouse and human obesity genes. For 4 of the 5, *Ksr2*, *Adcy3*, *Mc4r* and *Mc3r*, the KO mice are among the most obese we studied, with a 64% or greater increase in %BF relative to WT littermates, suggesting that the fidelity between mouse and human obesity genes may extend to the relative strength of their obese phenotypes. If so, then *GPR45*, *ASNSD1* and *TLE4* are the most likely of the remaining 65 genes to be confirmed as human obesity genes. Nevertheless, identification of obesity in individuals heterozygous for inactivating *NCOA1* mutations¹²⁰ despite the modest obesity of published^{119,120} and our *Ncoa1* KO mice suggests that homozygous inactivating mutations in human orthologs of any of the other 65 genes may be associated with severe human obesity. Despite the large effect of such homozygous inactivating mutations, their rarity precludes a major contribution to the polygenic obesity that results from small effects of many common genetic variants.¹⁶ However, homozygosity of complete loss-of-function mutations occurring simultaneously for 2 or more of these genes in the same person, undoubtedly a rare event, could result in morbid obesity due to the strength of the individual effects. This would most likely occur if the mutations in the 2 genes induce obesity through non-redundant pathways, resulting in an additive effect on fat mass; a good example is the increased body fat of *Mc3r*/*Mc4r* DKO mice, which equals the sum of the increased

Table 9 Shared Mouse Phenotypes for 18 Established Human Monogenic Obesity Genes

Human Gene	OMIM Gene	OMIM Disease	Shared Human and Mouse Phenotypes in Addition to Obesity and Hyperphagia	References	
				Human	Mouse
<i>ADCY3</i>	600291	None	Impaired glucose homeostasis; HET phenotype	[117,118]	[114,115]
<i>BDNF</i>	113505	None	Impaired glucose homeostasis; hyperactivity; HET phenotype	[216]	[241,242]
<i>CARTPT</i>	601606	601665	HET phenotype	[225]	[253–255]
<i>CEP19</i>	615586	615703	Impaired glucose homeostasis; insulin resistance; dyslipidemia; male infertility	[224]	[224]
<i>CPE</i>	114855	None	Impaired glucose homeostasis; hyperproinsulinemia; hypogonadotropic hypogonadism; impaired prohormone processing	[222]	[249–251]
<i>KSR2</i>	610737	None	Impaired glucose homeostasis; HET phenotype	[40]	[36,40,49,100]
<i>LEP</i>	164160	614962	Delayed puberty; mild hypothyroidism; insulin resistance; dyslipidemia; impaired immunity	[204–206]	[10,11,234]
<i>LEPR</i>	601007	614963	Delayed puberty; mild hypothyroidism; insulin resistance; dyslipidemia; impaired immunity	[207,208]	[10,12,234]
<i>MC3R</i>	155540	602025	Low lean body mass	[226,227]	[36,60–62,256]
<i>MC4R</i>	155541	618406	Impaired glucose homeostasis; increased linear growth and lean body mass; HET phenotype	[209–211]	[13,36,59,60]
<i>MRAP2</i>	615410	None	None	[220,221]	[220,248]
<i>NCOA1</i>	602691	None	HET phenotype	[120]	[119,120]
<i>NTRK2</i>	600456	613886	HET phenotype	[217]	[243,244]
<i>PCSK1</i>	162150	600955	Impaired prohormone processing; diarrhea; hyperproinsulinemia; HET phenotype	[228–233]	[232,257–261]
<i>POMC</i>	176830	609734	Defective adrenal function; altered pigmentation	[212,213]	[235–237]
<i>SH2B1</i>	608937	None	Impaired glucose homeostasis; maladaptive, aggressive behavior; HET phenotype	[214,215]	[238–240]
<i>SIM1</i>	603128	None	Increased linear growth; HET phenotype	[218,219]	[245–247]
<i>TUB</i>	601197	616188	Retinal dystrophy	[223]	[249,252]

Abbreviation: HET, heterozygous.

body fat found in the individual Mc3r and Mc4r KO lines ([Supplementary Table 4](#)).^{60,62}

The Value in Identifying Genes That Regulate Body Fat

The major value in identifying novel mouse obesity genes is that the product of the orthologous human gene is likely to regulate body fat content in man. One advantage of this conservation across species is that identifying a novel mouse obesity gene may suggest that, among many genes associated with a human GWAS BMI cluster, the

orthologous human gene is likely responsible for the signal. In support of this hypothesis, 10 of the 13 human obesity genes originally identified in mice are surrounded by or overlap with a GWAS BMI cluster (*LEP*, *LEPR*, *MC4R*, *POMC*, *SH2B1*, *BDNF*, *NTRK2*, *KSR2*, *ADCY3* and *NCOA1*), while the other 3 (*CPE*, *TUB* and *MC3R*) are located within 1 MBP of a cluster (<https://hugeamp.org/>). Thus, going forward, novel mouse obesity genes can focus research efforts on their orthologs located in poorly characterized BMI clusters. For example, a BMI cluster located within the *PRKDI* gene and 0.49 MBP 5' to the

G2E3 gene led to characterization of *PRKD1* as an obesity gene²⁶² and inclusion of *PRKD1* in a functional protein interaction network for childhood BMI²⁶³ despite the lack of functional data linking *PRKD1* to obesity. Our finding of obesity in *G2e3* KO mice but not in *Prkd1* KO mice⁴⁷ suggests that future work should focus on the role of *G2E3*, rather than *PRKD1*, in human obesity.

Knowing the gene mutation responsible for obesity in an individual may predict obesity risk in current and future family members and guide management of the biological effects associated with inactivation of that specific gene.^{264,265} In addition, some single gene mutations that result in obesity are associated with specific effective therapies. For example, recombinant leptin effectively treats obesity in people lacking functional leptin.²⁰⁶ Also, setmelanotide is a subcutaneously administered peptide, which decreases appetite and increases energy expenditure by selectively activating the MC4R through a novel and incompletely understood mechanism that avoids the cardiovascular toxicity of earlier generation MC4R agonists.^{266,267} Setmelanotide can effectively treat obesity in individuals with inactivating mutations in *LEPR*, *POMC* and *PCSK1* genes, which all encode proteins acting upstream of MC4R,^{268,269} and may also prove to be effective in individuals with specific *MC4R* mutations or with inactivating mutations in other genes, including *CPE* and *ADCY3*, that may result in impaired MC4R pathway function.^{266,268} This evidence supports the current recommendation for genetic testing of individuals with extreme early onset obesity, before the age of 5, that is associated with extreme hyperphagia, other features of genetic obesity syndromes, and/or a family history of extreme obesity.²⁶⁴

In humans with syndromic obesity, where obesity is one of multiple characteristic findings that constitute the syndrome, mouse KO models may also predict genes contributing to the obesity and suggest potential treatments. For example, *MAGEL2* is one of 4 paternally expressed genes located in a chromosome 15 locus associated with Prader-Willi syndrome or PWS (OMIM 176270). Among the characteristic findings of PWS is early failure to thrive followed by overeating and marked childhood obesity. *Magel2* KO mice reproduce many of the characteristic findings in PWS including obesity and hyperphagia,²⁷⁰ observations that preceded comparable findings in humans with truncating *MAGEL2* mutations.²⁷¹ *Magel2* KO mice may potentially have a role to play in the development of therapeutics that reverse the hyperphagia

and obesity of individuals with PWS or isolated *MAGEL2* deficiency.²⁷²

Perhaps the greatest value in identifying obesity genes is the potential that the gene product is a target for a therapeutic, or a therapeutic itself, that can lower body fat in people with common polygenic obesity. Retrospective analyses found that mouse KO phenotypes for most genes targeted by drugs correlate well with the effects of those drugs.^{29–31} These data are supported by 1) Lexicon's prospective development of neutralizing antibodies that reproduce in WT mice the phenotypes found in *Angptl4*,²⁷³ *Fzd4*,²⁷⁴ *Dkk1*,³⁷ *Notum*^{37,275} and *Angptl3*²⁷⁶ KO mice; 2) the prospective development by Regeneron Pharmaceuticals of a human therapeutic antibody that reproduces in humans the *Angptl3* KO phenotype;^{277,278} and 3) Lexicon's prospective development of small molecules that reproduce in rodents the phenotypes observed in *Sglt1*,^{39,279,280} *Notum*,^{28,275,281,282} *Aak1*,²⁸³ *Limk2*,²⁸⁴ *Rock1/2*²⁸⁵ and *Sgpl1*^{286,287} KO mice, along with small molecules that reproduce in multiple species including humans the *Sglt2* KO phenotype.^{39,288–291}

The above work confirms the ability of mouse KO phenotypes to model drug effects. Importantly, mouse KO phenotypes model what happens when a drug target is inhibited; thus, when inactivating a gene leads to obesity, drugs must activate the protein product of that gene, or the protein product must itself be an agonist. Agonist drugs are often developed from secreted protein and GPCR families, and both families have been mined for possible obesity therapies.^{23–25,31} The secreted protein leptin is, of course, a *LEPR* agonist, and leptin treatment of adults with leptin deficiency dramatically lowered their BW, body fat and food intake.^{205,206,292} However, treating common polygenic obesity with leptin achieved only modest weight loss,^{293,294} consistent with studies in rodents^{295–297} and inconsistent with broad use of leptin as a weight loss therapy. Also, agonists against the MC4R and HTR2C GPCRs have been developed. As noted above, while early MC4R agonists showed significant toxicity, the more recently developed agonist setmelanotide is less toxic and treatment for 4 weeks in 5 subjects with common polygenic obesity led to weight loss of ~0.9 kg/week; a better understanding of how setmelanotide works may lead to MC4R agonists that are safe and effective weight loss drugs for individuals with common polygenic obesity.^{266,298} Of more relevance here, the obesity of *Htr2c* KO mice⁶³ inspired development of the HTR2C agonist lorcaserin; *HTR2C* is neither associated with

a GWAS BMI cluster (<https://hugeamp.org/>) nor a human monogenic obesity gene. Lorcaserin, which has the advantage over peptide drugs of being an orally available small molecule, was approved by the FDA after demonstrating 5% weight loss in 49% of people with common polygenic overweight or obesity after 1 year of treatment.^{66,299–301} Lorcaserin was removed from the market in 2020 due to increased cancer risk, but development and approval of lorcaserin serves as a precedent for screening KO mouse models to identify obesity drug targets. Finally, identifying drug targets is not the only way that KO mouse models can aid obesity drug discovery. First, our KOs are global, they inactivate the targeted gene throughout the body, just as would a drug targeting the product of that gene; thus, thoroughly characterizing global KO mice may identify undesirable phenotypes that portend undesirable side effects of an on-target drug. Second, global KO models can be used to confirm that the anti-obesity effects of a drug are on-target, as shown by the ability of both lorcaserin and setmelanotide to lower food intake in WT mice but not *Htr2c* KO or *Mc4r* KO mice, respectively.^{65,302}

Confirmation of Novel Obesity Phenotypes in Mouse KO Models

The lack of reproducibility in published research has long been recognized but became a major issue a decade ago with reports that industry scientists could not reproduce most published preclinical studies.^{303–305} A recent update reviewed progress and emphasized that the research culture, particularly at academic institutions, remains a trans-cending challenge.³⁰⁶ These observations demand confirmation of all novel obesity phenotypes in mouse KO models; data from one KO model at one center is insufficient regardless of investigator, center, study design, data or *P* value. Confirmation of an obese mouse KO model can be based on studies of:

1. A KO of the same gene generated using an independent strategy by a different research group. This is the most frequent approach, and 43 examples are provided above.
2. A KO of the same gene generated using an independent strategy by the same research group. For example, Lexicon confirmed the obesity phenotype of *Ksr2* KO mice by reproducing the phenotype in an independent *Ksr2* KO model.⁴⁹
3. A KO of a different gene that expresses the sole ligand or receptor of the gene product of the obese KO model. *Lep/Lepr* (ob/ob and db/db) and α -MSH and *Mc4r* (*Pomc/Mc4r*) are examples that support this approach.^{10–13,235} As examples from this report, we used our obese *Kiss1r* KO mice to first confirm the published phenotype^{69,70} and then to confirm the obesity of our *Kiss1* KO mice, which lack the ligand for KISS1R; in addition, the strikingly similar obesity phenotypes of *Prok2* KO and *Prok2r* KO mice suggest roles for each gene in regulating body fat.
4. A KO or inactivating mutation of the same gene in a different species. The best examples are mouse KOs for *Mrap2*,²²⁰ *Cep19*²²⁴ and *Cartpt*,^{225,253} which were simultaneously reported as human obesity genes, and KOs for *Sim1*^{218,245} and *Pcskl*,^{228,257–261} which were first reported as human obesity genes. In addition, human GWAS data may provide support for a KO obesity phenotype; for example, our *Tle4* KO mice are obese and a human GWAS BMI cluster sits within the *TLE4* gene, which is at the center of, and the only gene in, a 2.5 MBP stretch of human chromosome 9.
5. The effect of a drug that specifically targets the gene product of the obese KO model. The best example is the HTR2C agonist lorcaserin, which lowers body fat of WT mice but not *Htr2c* KO mice.⁶⁵

Often, multiple approaches are combined to confirm that a gene is linked to obesity. For *Ksr2*, confirmation was achieved by observing 1) obesity in two independent in-house KO models; 2) obesity in an external KO model; 3) a strong link of the human *KSR2* gene to GWAS BMI clusters; and 4) obesity in humans with *KSR2* haploinsufficiency.^{36,40,49,100} Rather than depend on data from one KO cohort, confirmation requires supporting data from multiple cohorts, following the Bayesian statistical paradigm.³⁰⁷ In the absence of supporting data, there is a real risk that the obesity is a false-positive observation. This caution is particularly appropriate for KO lines that are linked to obesity based exclusively on HTS data. The obesity of these KO lines is best viewed as a hypothesis-generating observation requiring confirmation using an independent KO model.

If a mouse KO obesity phenotype is not confirmed with an independent mouse KO model, there are many issues to consider that may explain the discrepancy:

1. Study power.³⁰³ Study power depends on strength of phenotype and the number of mice studied/group. As shown in Table 1, the strength of obesity phenotypes characterized under very similar conditions has a wide range, with the well-established *Mc4r* KO and *Prhr* KO phenotypes differing greatly in strength. Power calculations can determine if the number of mice/group was adequate to observe the phenotype in the original and/or confirmatory studies and are particularly important to perform for subtle phenotypes.
2. Bias.³⁰³ Study methods and interpretation should be reviewed for evidence of bias. Agnostic approaches, such as our HTS approach, are one way to minimize the contribution of bias to study results.
3. Mouse KO strategy. Homozygous inactivating *PCSK1* mutations led to obesity in humans but not mice; however, obesity was observed in more targeted mouse *Pcsk1* mutations and in HET mice.^{228,257–261} Also, an initial *G2e3* KO strategy resulted in embryonic lethality, but two recent and different KO strategies both found that *G2e3* KO mice were obese.⁴⁷ Of note, in our experience gene trap and homologous recombination technologies produced comparable KO phenotypes for *Angptl4*,²⁷³ *Tph1*,³⁰⁸ and *Adams4* (this manuscript), suggesting that differences between KOs generated by these two techniques are more likely due to differences in specific mutation strategies rather than differences in the two techniques.
4. Mouse phenotyping strategy. In our experience, body fat data derived from DXA and QMR technologies are comparable and accurate surrogate measurements of body fat; excellent correlations were found between % body fat measured by both QMR and DXA and by both QMR and carcass analysis.³⁶ However, BW is a less sensitive surrogate measurement for body fat, as exemplified by multiple cohorts of chow fed *G2e3* KO and WT mice that showed significant differences in body fat measured by QMR but not in BW.⁴⁷ For many KO lines, a subtle but significant increase in body fat detected by DXA and/or QMR was not accompanied by a significant increase in BW.
5. Mouse age. Obesity would be missed in many mouse KO lines if body fat was measured only at weaning; for example, *Ksr2* KO and *Adcy3* KO mice show reduced viability with low BW and body fat at weaning but rapidly develop obesity over the next few weeks,^{49,114} similar to our findings here for *Tle4* KO mice.
6. Mouse genetic background. Classic studies show that the obesity of *ob/ob* (*Lep*) and *db/db* (*Lepr*) mice is influenced by background mouse strain.^{10,309} Similarly, cold tolerance is decreased when *Ucp1* KO mice are on a C57BL/6J or 129/SvImJ background but not when on a hybrid C57BL/6J X 129/SvImJ background.³¹⁰ Background may also explain why *Htr2c* KO mice were obese on a C57BL/6J background^{63,64} but not on our hybrid C57BL/6J X 129SvEvBrd background. We often saw phenotypes in adult KO mice maintained on our hybrid background when embryonic lethality was reported for an IMPC KO of the same gene maintained on their C57BL/6N background;²⁸ this hybrid vigor is likely due to a different and broader complement of modifier genes in our KO lines that allows survival and phenotyping.
7. Mouse diet. Mouse diet likely influences the strength of the obesity phenotype for certain mouse KO lines. For example, *Pnpla2* KO mice were reported as obese when fed chow diet but not HFD¹³⁵ and *Ddahl* KO mice were reported as obese when fed HFD but not chow diet,¹³⁸ observations supported by our data for each KO line. However, for almost all KO lines reported here, obesity was observed regardless of whether KO and WT littermate mice were fed chow or HFD.
8. Mouse sex. Sexual dimorphism has been reported for many phenotypes but was rare in our skeletal³⁷ and obesity phenotyping programs; in our experience, sex differences in either bone or fat HTS data were usually spurious findings associated with an overall weak phenotype that was not reproduced when additional cohorts were studied. Confirming the finding of obesity in mice of a single sex requires additional adequately powered studies that provide data on both male and female KO mice.
9. Environment. Environment can influence the amount of body fat stored. For example, *Ucp1* KO mice were more likely to be obese at thermoneutrality than at lower temperatures³¹¹ and *Mc4r* KO mice were obese in the absence, but not the presence, of running wheels.³¹² These observations suggest that researchers evaluate the effects of specific environmental conditions on the phenotype

they are studying; ambient temperature, cage size and numbers of mice/cage can all influence body fat accumulation.

10. Use of large-scale mouse KO HTS data for confirmation. The ongoing IMPC and the completed Lexicon Pharmaceuticals Genome5000TM programs are the two large-scale mouse phenotyping campaigns that used reverse genetics, which analyzes phenotypes resulting from KO of specific genes, to provide data on the physiologic function of those genes. Their major advantage is using a panel of assays with standardized protocols to discover, in an unbiased manner, a broad range of phenotypes in mice. Their major disadvantages are that mouse cohorts are small and are studied within a narrow age range, creating a risk for false positive and negative results. These and other issues are discussed in a recent review of these two campaigns²⁸ but a few observations on the fidelity of phenotyping by the Lexicon HTS are warranted. Importantly, each KO line compares KO and WT littermates/cagemates for each assay, which controls for environmental differences and for genetic drift of the colony over time. The Lexicon HTS identified established, strong body fat and skeletal KO phenotypes with high fidelity, including correct identification of the body fat phenotype of all 5 KO lines with established published obesity.^{28,36,37} Novel subtle phenotypes were more challenging. One strength of the Lexicon HTS was that 2 independent cohorts were usually screened for body fat, one fed chow and the other HFD from weaning, which provided insight into the consistency of a body fat phenotype. Nevertheless, the modest obesity observed by HTS in some KO lines was not reproduced when additional in-house cohorts were studied even when both chow- and HFD-fed HTS cohorts appeared obese.³⁶ False-positive phenotypes are also expected for IMPC HTS KO lines.³¹³ Of the 29 KO lines reported here with the novel finding of increased body fat in HTS KO mice, body fat data for 2 of the KO lines (*Ksr2* and *G2e3*) were supported by data from additional internal and external KO cohorts and for 6 of the KO lines were supported by data from additional internal cohorts, but body fat data for the remaining 21 KO lines were unsupported by data from additional cohorts and are at higher risk of being false-positive observations. Ultimately, the novel finding of high body fat in 27 of our KO lines

is a hypothesis-generating observation for each line that requires demonstration of significant obesity in an independent external KO model for confirmation.

Summary

Lexicon Pharmaceuticals used high-throughput approaches to both create KO lines for mouse genes that had drug-gable human orthologs and to analyze body fat in adult littermate/cagemate WT and KO mice from each KO line. Body fat was usually analyzed in 2 independent cohorts of these mice, one maintained on chow diet and the other on HFD from weaning. This program identified dozens of obese KO lines; for many of these lines, the obesity observed in the KO mice confirmed data from previously established KO models, while for others the obesity was a novel finding that will require external confirmation. Undoubtedly, many mammalian obesity genes remain to be identified and characterized.

Acknowledgments

The authors wish to thank all Lexicon animal care, phenotyping and IT professionals who worked diligently to generate and catalog the data presented here, Kristi Boehm, MS, ELS, for her help in editing and in preparing the figures and tables, and Dr. Sadia Saeed for helpful discussions.

Disclosure

All authors were employed by Lexicon Pharmaceuticals, Inc., at the time these studies were performed and may own common stock or may have been granted stock options or other equity incentive awards. The authors report no other conflicts of interest in this work.

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