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Cre/lox generation of a novel whole-body *Kiss1r* KO mouse line recapitulates a hypogonadal, obese, and metabolically-impaired phenotype

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Abstract

Kisspeptin and its receptor, *Kiss1r*, act centrally to stimulate reproduction. Recent evidence indicates that kisspeptin is also important for body weight and metabolism, as whole-body *Kiss1r* KO mice, developed with gene trap technology, display obesity and reduced metabolism. *Kiss1r* is expressed in brain and multiple peripheral tissues, but it is unknown which is responsible for the metabolic phenotype. Here, we sought to confirm that 1) the metabolic phenotype of the gene trap *Kiss1r* KOs is due to disruption of kisspeptin signaling and not off-target effects of viral mutagenesis, and 2) the *Kiss1r* flox line is suitable for creating conditional KOs to study the metabolic phenotype. We used Cre/lox technology (*Zp3-Cre/Kiss1r* flox) to develop a new global *Kiss1r* KO ("*Kiss1r* gKO") to compare with the original gene trap KO phenotype. We confirmed that deleting exon 2 of *Kiss1r* from the entire body induces hypogonadism in both sexes. Moreover, global deletion of *Kiss1r* induced obesity in females, but not males, along with increased adiposity and impaired glucose tolerance, similar to the gene trap *Kiss1r* KOs. Likewise, *Kiss1r* gKO females had decreased VO_2 and VCO_2 , likely underlying their obesity. These findings support that our previous results in gene trap *Kiss1r* KOs are due to disrupted kisspeptin signaling, and further highlight a role for *Kiss1r* signaling in energy expenditure and metabolism besides controlling reproduction. Moreover, given *Kiss1r* expression in multiple cell-types, our findings indicate that the *Kiss1r* flox line is viable for future investigations to isolate specific target cells of kisspeptin's metabolic effects.

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Keywords

kisspeptin; Kiss1; Kiss1r; GPR54; adipose; fat; obesity; metabolism; energy expenditure

Introduction

Kisspeptin (encoded by the gene *Kiss1*) is a neuropeptide that binds the receptor, Kiss1r (previously termed GPR54). In all mammals, including humans, kisspeptin and its receptor regulate puberty and reproduction by acting in the brain to directly stimulate GnRH secretion, evidenced by findings that humans and rodents with disrupted kisspeptin signaling are infertile, have undeveloped gonads (hypogonadal), and have severely diminished reproductive hormone secretion (reviewed in (1–5)). In addition to being expressed in GnRH neurons in the brain (6–8), *Kiss1r* is also located in several peripheral tissues, such as adipose tissue, pancreas, adrenal, and gonads (9–14). Likewise, kisspeptin is expressed in several peripheral tissues, including liver, pancreas, gonad, and placenta (12–14). This peripheral expression of both kisspeptin and its receptor suggests that kisspeptin signaling has additional roles besides reproductive regulation. However, until recently, non-reproductive roles of kisspeptin have been largely overlooked.

Our group recently reported that, in addition to stimulating the reproductive axis, kisspeptin signaling also influences body weight (BW), energy balance, and glucose homeostasis. Specifically, compared with WT littermates, global *Kiss1r* knockout (KO) female mice—lacking kisspeptin signaling in all tissues—displayed significantly greater BWs in adulthood, weighing as much as 30% more than control females (15,16). In addition to becoming obese, *Kiss1r* KO females also had increased adiposity, higher leptin levels, and substantially impaired glucose tolerance, both on standard chow and high fat diets (15,16). This obesity phenotype was sexually dimorphic, with *Kiss1r* KO males exhibiting normal BW and glucose regulation out to 6 months of age. The obese *Kiss1r* KO females did not have increased food intake, but rather they displayed significantly reduced metabolism parameters, including highly reduced metabolic respiratory rates and energy expenditure. These data suggests that their obesity reflects lower daily energy expenditure (15,16) rather than changes in food intake. Importantly, the obesity and metabolic dysfunction in *Kiss1r* KO females was not just due to their hypogonadal state and absent ovarian estradiol (E₂), as long-term ovariectomized *Kiss1r* KO females still developed obesity, hyperleptinemia, reduced metabolism, and glucose intolerance when compared with long-term ovariectomized WT control females (15,16). In both genotypes, E₂ was absent for many month starting from well before puberty, yet the KOs still became heavier and had lower metabolic rates, signifying the phenotype is not just caused by their hypogonadal state.

Because Kiss1r was completely absent from all cells in the body in the global *Kiss1r* KOs, it remains unknown which specific tissues and cell types, neural or peripheral, are responsible for kisspeptin's influence on the various metabolic phenotypes. Indeed, peripheral kisspeptin and Kiss1r are present in several metabolic-related peripheral tissues, suggesting a number of potential candidate target sites for kisspeptin's ability to alter metabolism and energy balance. Moreover, it is possible that different phenotypic impairments (glucose homeostasis

impairment, increased adiposity, altered metabolic rate, etc.) observed in global *Kiss1r* KOs may each reflect absent kisspeptin signaling in different tissues or cell-types, with the net effect of each of these individual tissue-specific changes resulting in the overall obese phenotype. To dissect and isolate the contribution of specific individual cell types/tissues to the various metabolic alterations, future studies might utilize Cre-driven selective knockout of kisspeptin signaling from just one tissue at a time and compare this to the condition in whole-body global KO females. To date, this has not been attempted, and first requires validation that available *Kiss1r* floxed mouse lines are able to recapitulate the global KO phenotype when used to generate full body KOs.

Multiple strategies have been used to generate mutations in rodent models, including gene targeting by homologous recombination, point mutations or small deletions generated by chemical mutagenesis, and random gene trap mutagenesis applied genome-wide using plasmid, virus, or transposon DNA vectors delivered to ES cells (17–19). Our global *Kiss1r* KO mouse displaying the obese, metabolic phenotype was created via gene trapping by retroviral mutagenesis. When inserted into an intron of an expressed gene, the gene trap cassette causes transcription to be terminated prematurely, resulting in a fusion transcript encoding a truncated, nonfunctional protein (17,20). Despite other known *Kiss1r* and *Kiss1* KO mouse lines similarly having a hypogonadal, infertile phenotype, confirmation of a metabolic/obese phenotype in these other KO lines has yet to be reported. It is unclear if this is because metabolic parameters have not yet been extensively studied in the other KO lines or if our retroviral gene-trap variant exhibits a metabolic phenotype that is not recapitulated by other genetic models of kisspeptin impairment.

The present study experimentally tested whether a novel whole-body KO line created with Cre/lox technology could replicate the previous findings in our original global “gene trap” *Kiss1r* KO line, thereby providing further evidence for a metabolic role of kisspeptin signaling. By crossing our recently-made *Kiss1r* flox line (21) with *zona pellucida 3-Cre* (*Zp3-Cre*) mice, we generated a new total body deletion of *Kiss1r*. The *Zp3-Cre* mouse line utilizes Cre expression controlled by regulatory sequences from the mouse *Zp3* gene, which is expressed exclusively in the growing oocyte prior to the completion of the first meiotic division (22–25). First, we used this novel *Zp3-Cre/Kiss1r* flox model of whole body *Kiss1r* deletion to validate and confirm of our previous report of obesity and metabolic disturbance in global *Kiss1r* KO mice made using retroviral gene trap technology. Secondly, from a conceptual and technical perspective, we used this novel approach to provide essential proof of feasibility for future Cre/lox studies that aim to selectively knock out *Kiss1r* in just one specific metabolic cell type or tissue, in order to isolate and study kisspeptin’s site(s) of action underlying the obese phenotype. Herein we report the presence of a full metabolic, obese phenotype, along with severe hypogonadism, in these novel global *Kiss1r* KO mice generated with Cre/lox technology.

Materials and Methods

Animals

Experiments used a new global (whole-body) *Kiss1r* KO mouse line generated with Cre/lox technology to ascertain if one could recapitulate the previously-observed reproductive and

metabolic phenotypes of our original global *Kiss1r* KO mice that were created by gene-trap methodology (15,16,26,27). To create the new global KO of *Kiss1r*, *Zp3-Cre* female mice (24) were mated with males of a recently created *Kiss1r* flox line (*Kiss1r^{fl/fl}*) (13,21), allowing recombination of the *Kiss1r* allele to occur in the oocyte of female offspring. In target-bearing *Zp3-Cre* mice, Cre-mediated recombination of the target gene occurs in 100% of oocytes, and Cre expression is directed to the time when maternal genes are first transcribed, thereby inactivating target floxed genes in the maturing oocyte. This allowed us to effectively induce germline recombination to delete the target gene, *Kiss1r*, from the entire embryo (i.e., a global KO). We then selected recombination positive, Cre negative (*Zp3-Cre^{-/-}/Kiss1r^{rec/wt}*) offspring to further generate global *Kiss1r* KOs lacking functional *Kiss1r* in all cells (*Zp3-Cre^{-/-}/Kiss1r^{rec/rec}*; termed “*Kiss1r* gKO”) and control littermate mice (*Zp3-Cre^{-/-}/Kiss1r^{rec/wt}* and *Zp3-Cre^{-/-}/Kiss1r^{wt/wt}*).

All mice were genotyped and sexed by using PCR of DNA obtained from toe samples at postnatal day 7 (PND7) or tail DNA at weaning (PND 21). Weaned mice were housed 2–3 per cage with mixed genotype in a 12-hour light/12-hour dark cycle, with *ad libitum* water and standard rodent chow containing 3.5 kcal/g, 45.2% carbohydrates, 11.4% fat, and 17.2% crude protein. All experiments were approved by the Institutional Animal Care and Use Committee from the University of California San Diego.

Reproductive Characteristics

Previously reported global *Kiss1r* KO mice are well-known to have impaired reproductive status, as evidenced by extremely underdeveloped gonads, reduced anogenital distance (AGD; an indirect measure of testosterone exposure) in males, and correspondingly low circulating estrogen and testosterone levels. For the new global *Kiss1r* KO mouse line here (*Kiss1r* gKO), reproductive developmental status was quantified by measuring AGD at week 12 and gonadal weights at sacrifice. For the former, mice were briefly anesthetized with isoflurane and distance measurement was taken from the anus opening to the base of the reproductive organ.

Body Weight Assessment

To assess whether the novel *Kiss1r* gKOs develop a metabolic phenotype, as occurs in the gene trap global KOs, *Kiss1r* gKOs of both sexes were analyzed for their BW at multiple ages. Control and experimental littermates were weighed once every two wks, starting at 4 wks of age and ending around 4–5 months old.

Metabolic and Energy Expenditure Analyses

To measure metabolic rates, *Kiss1r* gKO and control littermates were analyzed at 20–24 weeks of age with indirect calorimetry using equal flow Comprehensive Laboratory Animal Monitoring System (CLAMS) calorimeter system (CLAMS; Columbus Instruments, Columbus, OH). Mice were individually housed 2–3 days prior to the experiment for habituation to being single housed. CLAMS data were collected in clear respiratory chambers (20 × 10 × 12.5 cm) equipped with a sipper tube delivering water and a food tray connected to a balance. The consumption of O₂ and production of CO₂ was measured by having sample air sequentially passed through O₂ and CO₂ sensors (Columbus Instruments)

for determination of O₂ and CO₂ content, from which measures of oxygen consumption (VO₂) and carbon dioxide production (VCO₂) were estimated. Room air was passed through chambers at a flow rate of 0.5 L/min. Exhaust air from each chamber was sampled at 15-min intervals for 1 min. Outdoor air reference values were sampled after every 8 measurements. Gas sensors were calibrated prior to the onset of experiments with primary gas standards containing known concentrations of O₂, CO₂, and N₂ (Airgas Puritan Medical, Ontario, CA). Respiratory exchange ratio (RER) was calculated as the ratio of carbon dioxide production (VCO₂) to oxygen consumption (VO₂). Energy expenditure (heat formation) was corrected for each mouse's body mass. Daily food intake was also recorded by the CLAMS system in a subset of mice of each genotype. All data were recorded under ambient room temperature (~24°C) for up to 7 days.

Body Composition Analyses

Lean and fat mass of a group of *Kiss1r* gKO females and males was determined using dual energy x-ray absorptiometry (DEXA) at 19–21 weeks old. The mice were fasted for 4–6 hours before measurement. The mice were anesthetized by an ip injection of Fatal-Plus. The DEXA scan was performed with a GE Lunar Pixi Densitometer Machine.

Glucose Tolerance Tests

At 18–20 weeks of age, *Kiss1r* gKO mice and their control littermates of both sexes underwent glucose tolerance testing (GTT). Mice of both genotypes and sexes were fasted for 6 hours before starting the glucose measurements, with free access to water throughout the experiment. Blood glucose levels were measured using a handheld glucometer before (time 0 min) and after (15, 30, 45, 60, 90, and 120 min) a single ip glucose injection (2g/kg BW dissolved in saline).

Assessment of *Kiss1* and *Kiss1r* gene expression levels in both sexes

qRT-PCR analyses of *Kiss1* and *Kiss1r* genes was compared between normal (i.e., not KO) adult males and females (8 wks old) for multiple metabolically-relevant tissues, including whole hypothalamus, liver, white and brown adipose (WAT and BAT), and skeletal muscle. Both sexes were GDX 2 wks prior to tissue collection to equalize circulating sex steroids between all the animals. Total RNA was isolated using the phenol/chloroform method and converted to cDNA via reverse transcription (Promega, Sydney, Australia), followed by clean up using the PCR Clean Up Kit (Mo Bio Laboratories, Carlsbad, CA, USA). qRT-PCR was performed in 10- μ l reaction volumes and samples were tested in duplicate using a Rotorgene 3000 (Corbett Life Science; Qiagen, Valencia, CA, USA). Reactions used Qiagen SYBR Green PCR Kit (Qiagen, Chadstone Centre, VIC, Australia). Samples were compared with a standard curve (10-fold dilution) and relative gene expression was normalised using a GE Norm algorithm of housekeeping genes peptidylpropyl isomerase A (*Ppia*), succinate hydrogenase (*Sdha*) and TATA box binding protein (*Tbp*). No significant variability was noted in each housekeeping gene.

Statistical Analyses

All data are presented as mean \pm SEM. For data at single points (non-repeated measures), single comparisons were made using one- or two-tailed t-tests, as appropriate, and multiple comparisons were performed using one-way ANOVA with Tukey's post-hoc test. For repeated measures (BWs, GTTs), repeated measures ANOVA was performed, with Bonferroni post-hoc tests directly comparing genotypes at specific points for 3 groups or t-tests for 2 groups. Statistical significance was $p < 0.05$.

Results

Reproductive development is impaired in a new global *Kiss1r* KO line generated with Cre/lox technology

We sought to 1) confirm that disrupting global *Kiss1r* signaling using Cre/lox technology similarly caused the metabolic phenotypes previously observed in our original gene trap global *Kiss1r* KOs, and 2) validate that the new *Kiss1r* flox line is a viable tool for future testing of site-specific kisspeptin metabolic actions. We therefore generated a new global *Kiss1r* KO line ("*Kiss1r* gKO") via germline Cre using *Zp3*-Cre mice mated to *Kiss1r* flox mice to delete the second exon of *Kiss1r* in all cell types throughout the body (Figure 1A and 1B). Given kisspeptin's established role in promoting reproductive development and fertility, we first assessed gonadal status of these new *Kiss1r* gKO mice. Adult *Kiss1r* gKO of both sexes had significantly smaller, undeveloped gonads (testes in males and ovaries in females) compared to WT controls ($p < 0.001$; Figure 1C), indicating that the reproductive axis was impaired, as expected, in our new global *Kiss1r* KO line. Supporting this, AGD (a testosterone-stimulated somatic measure) in adult *Kiss1r* gKO males was significantly smaller than in WT male littermates ($p < 0.01$; Figure 1D).

Body weight are elevated in the new global *Kiss1r* KO line

To determine if the new *Kiss1r* gKOs develop an obesity phenotype like our previous gene trap global *Kiss1r* KO females, BWs were measured every two weeks starting at week 4 (before puberty). Adult *Kiss1r* gKO females showed a significantly higher BW compared to WT female littermates ($p < 0.01$; Figure 2A), starting around week 12 and continuing through adulthood. By week 18 of age, *Kiss1r* gKO females weighed a marked 30% more than WT littermate controls. Conversely, *Kiss1r* gKO males did not display any BW differences compared to their WT littermates (Figure 2B), similar to the gene trap global KO males.

Adiposity is elevated in the new global *Kiss1r* KO line

Adult *Kiss1r* gKO and WT females were analyzed by DEXA to see if their body fat and lean mass composition were altered. gKO females showed a significantly higher fat mass and percent fat mass compared to WT females ($p < 0.001$ for each measure; Figure 3A and 3B), while their lean mass was significantly lower than WT females ($p < 0.01$; Figure 3C). Circulating leptin was also significantly elevated in female *Kiss1r* gKOs ($P < 0.01$; Figure 3D), matching their elevated fat mass. In line with their normal BWs, *Kiss1r* gKO males did not show any significant differences in fat mass compared to WT males (*data not shown*).

Glucose tolerance is impaired in the new global *Kiss1r* KOs created with Cre/lox technology

We performed an ip GTT to assess glucose regulation in adult female *Kiss1r* gKOs. There was a significant genotype difference, with *Kiss1r* gKOs displaying impaired glucose tolerance (i.e., higher blood glucose levels and slower clearance) relative to WT control females, similar to the gene trap global KOs ($p < 0.01$; Figure 4).

Metabolic rates are significantly reduced in the new global *Kiss1r* KO line

Despite their obesity, mean daily (24 h) food intake was significantly lower in the *Kiss1r* gKO females than in WT littermate controls (2.41 ± 0.02 vs 3.37 ± 0.08 g/day, $p < 0.01$). To test whether the obesity in the female *Kiss1r* gKOs may be due instead to diminished metabolism, we used CLAMS metabolic cages to measure metabolic respiratory rates. Adult *Kiss1r* gKO females had significantly lower levels of both O_2 consumption and CO_2 production during both the light and dark phases of the light-dark cycle relative to WT female littermates ($p < 0.01$; Figure 5A and B), similar to the gene trap global *Kiss1r* KO females. Similarly, adult *Kiss1r* gKO females had significantly lower energy expenditure during both the light and dark phases relative to WT females ($p < 0.5$; Figure 5C).

Expression patterns of *Kiss1* and *Kiss1r* in metabolic tissues from males and females

Our results above indicated a major sex difference in the metabolic effects of global *Kiss1r* deletion: both male and female *Kiss1r* gKO mice showed hypogonadism, but only female *Kiss1r* gKOs showed obesity and metabolic impairment. To try and understand the underlying cause of this metabolic sex difference, we compared gene expression levels of *Kiss1r*, as well as *Kiss1* (encoding the ligand, kisspeptin), in several metabolic tissues of male and female mice. In both sexes, both the receptor and ligand mRNAs were detected in the hypothalamus and several peripheral metabolic tissues (Figure 6), as previously reported. In the peripheral tissues, *Kiss1* was most highly expressed in liver whereas *Kiss1r* was expressed more abundantly in WAT and BAT. However, there were no notable sex differences in either *Kiss1r* or *Kiss1* gene expression in any tissue examined (Figure 6), suggesting that the obesity sex difference in gKOs reflects an underlying cause other than differential *Kiss1* or *Kiss1r* expression.

Discussion

Kisspeptin-Kiss1r signaling in the brain is widely acknowledged as a critical activator of the reproductive axis in all mammals, including humans. Kisspeptin's reproductive effects are achieved through its neural actions, stimulating GnRH neurons, which express Kiss1r. However, recent evidence from our group and others demonstrate that kisspeptin-Kiss1r signaling is also important for non-reproductive processes, including metabolism and energy balance. We recently reported that global *Kiss1r* KO females, generated via gene trap technology, develop obesity and glucose intolerance in adulthood, due to markedly decreased metabolic rates and energy expenditure that is independent of E_2 levels (15,16,26). However, because Kiss1r is expressed in several peripheral tissues, the specific tissue(s) responsible for the observed obese/metabolic phenotype in the global KOs remains undetermined. Here, we demonstrated that the previously-observed obese and metabolic

phenotype could be completely reproduced via Cre/lox technology by deleting exon 2 of *Kiss1r* in the oocyte, thereby producing germline recombination that targets all cells throughout the body (24). Thus, endogenous kisspeptin signaling somewhere, in one or more target cell-types, provides a previously-unrecognized modulation of energy expenditure, body weight, and metabolism.

Our prior studies found that our original gene trap global *Kiss1r* KO mice are hypogonadal and infertile, similar to other reported mouse knockout models of either *Kiss1* or *Kiss1r* (21,28–31). Our present results, using Cre/lox technology (*Zp3-Cre/Kiss1r* flox, aka *Kiss1r* gKO), confirmed that deleting the second exon of *Kiss1r* from the entire body results in hypogonadal female and male mice. *Kiss1r* gKO males also had reduced AGD, a testosterone-dependent measure. These findings indicate that Kiss1r signaling in reproductive cells (likely GnRH neurons (21,30)) was disrupted, as expected, validating the *Kiss1r* flox line. The Cre/lox methodology used here is therefore viable to generate *Kiss1r* KOs for future reproductive investigations.

Metabolic assessments in the newly generated *Kiss1r* gKO model also confirmed that lacking *Kiss1r* from the entire body results in obesity in females, but not males, with higher fat mass and lower lean mass, similar to the original global *Kiss1r* KO females. Similarly, *Kiss1r* gKO females also had significantly lower VO₂ and VCO₂ compared to control females, likely underlying the observed obesity in the *Kiss1r* gKOs. Daily food intake was not elevated in the *Kiss1r* gKOs, but rather was decreased relative to controls, as in the gene-trap global KOs. The lower food intake may be a side effect of the elevated leptin in these obese females, as leptin is known to diminish feeding (32–34), but this remains to be tested in future studies. All of these findings confirm that our previous results in the gene trap KO animals are due to disruptions in kisspeptin signaling, and not off target effects of viral mutagenesis. These findings also confirm that Kiss1r signaling is important for not only stimulating the reproductive axis, but also for regulating BW, body composition, and metabolism, at least in females. Importantly, in support of this, a recent clinical study in humans reported metabolic-related effects of kisspeptin (35). In that study, a single infusion of kisspeptin to patients significantly increased insulin secretion and also altered serum metabolites, including reducing small lipid species in the blood (35). These findings in humans are congruent and complementary with our present *Kiss1r* gKO findings and previous KO reports (15,16), which suggested that functional kisspeptin signaling may normally act to promote insulin action, glucose homeostasis, and prevent elevated adiposity. It will be interesting to see if future clinical studies report alterations in BW after longer-term kisspeptin treatment, and if kisspeptin treatment increases metabolic rates or body temperature in humans.

Given that Kiss1r and kisspeptin are both present in several peripheral metabolic tissues (Figure 6), the exact location(s) of where kisspeptin signaling influences these metabolic and energy balance parameters remains unknown. Of note, recent data indicate that neural energy balance populations, like hypothalamic neuropeptide Y (NPY) and pro-opiomelanocortin (POMC), have unaltered metabolic gene expression in global *Kiss1r* KO females (26), suggesting that the underlying mechanism giving rise to obesity and reduced metabolic rates in those mice is likely occurring outside the brain in peripheral systems. To

test this possibility, future studies can employ Cre/lox technology to conditionally delete *Kiss1r* specifically from a target cell type or tissue, while maintaining intact *Kiss1r* elsewhere in the body. The present study confirms that this approach is technically feasible, as the present *Kiss1r* flox line, when crossed with *Zp3-Cre* line, was able to be used to fully recapitulate the full metabolic phenotype observed in the global *Kiss1r* KO mice used in previous studies. In addition to the brain, *Kiss1r* is expressed in white and brown adipose tissue, liver, gonads, and pancreas, so determining the exact functional target site will require extensive examinations. A previous study found that deletion of *Kiss1r* from just the pancreas improved glucose metabolism, but that study did not measure body weight, adiposity, or metabolic rates and energy expenditure (13). Thus, it remains unknown if kisspeptin action in the pancreas contributes to BW and metabolic regulation in addition to possible effects on insulin secretion. Regardless, it is important to note that the observed metabolic phenotype of the *Kiss1r* KOs may not be due to absent kisspeptin signaling in just one tissue. Indeed, actions of kisspeptin in one specific tissue type may contribute to only one or two components of the “overall” metabolic phenotype whereas kisspeptin acting in other specific tissue type may underlie other metabolic aspects. For example, hypothetically, kisspeptin signaling in the pancreas may influence insulin and glucose tolerance, whereas kisspeptin signaling in adipose tissue may influence adiposity or energy expenditure.

Unlike the females, Cre/lox-derived *Kiss1r* gKO males did not display obesity or alterations in body fat, similar to what was observed in the gene trap global *Kiss1r* KO males. Despite the absence of a metabolic phenotype, the *Kiss1r* gKO males did display severe hypogonadism and reduced AGD, signifying that the Cre/lox deletion of *Kiss1r* signaling was successful. The mechanistic reason for the obesity sex difference similarly observed in both the original global KOs and the new *Kiss1r* gKOs remains unknown. We tested whether there were sex differences in *Kiss1r* or *Kiss1* expression levels in several tissues but found similar mRNA levels between males and females in all tissues examined. This suggests that the obesity sex difference is not due to sex differences in peripheral kisspeptin production or sex differences in *Kiss1r* expression in metabolic target tissues, though it remains possible that there are unknown sex differences in these genes in other tissues or cell-types that were not tested in our present study, such as pancreas or adrenals. Investigating this issue will require well-controlled experimental designs given that *Kiss1* or *Kiss1r* expression in the periphery may be influenced by metabolic status, hormones, or other factors (12,13,36–38), as it is in the brain. Alternatively, the obesity sex difference may reflect differences in intra-cellular signaling pathways downstream of *Kiss1r* in specific metabolic target sites, a possibility for future investigations.

Of note, a recent report (39) from Tena-Sempere’s group analyzed body weight and adiposity phenotype in a different global *Kiss1r* KO line. They showed lower body weight gain in their KOs of both sexes at early ages (< 6 wks old), which we also previously reported in global *Kiss1r* KO males (16). Importantly, at older ages, their *Kiss1r* KO males and females showed normal or increased BW gain, respectively, similar to our prior findings. Moreover, adiposity levels were higher in KOs of both sexes, and on a high fat diet, their female KOs became fatter than WT controls, also matching our prior findings (39). Thus, there was generally good congruence between their findings and both our previous and current report.

In summary, global deletion of *Kiss1r* from all tissues/cells achieved via Cre/lox technology, results in both impaired reproductive status and diminished energy balance and metabolism in females, the latter of which was correlated with increased adiposity and BW. This confirms our previous findings in gene trap derived global *Kiss1r* KO mice and further supports a role for Kiss1r signaling in energy expenditure and metabolism in addition to just governing reproduction, findings recently supported by clinical studies in humans. Overall, these findings highlight the complexity and multi-faceted nature of the kisspeptin system throughout the body, and emphasize the need for more tissue- or cell-specific assessments to compare with global whole-body manipulations. Future studies are needed to determine the underlying mechanisms and specific sites of kisspeptin action that lead to obesity and metabolic dysfunction in global KOs, as well as to further define the specific actions of endogenous kisspeptin signaling specifically in target metabolic cell-types.

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Highlights

- We used Cre/lox technology to develop a novel global *Kiss1r* knock out (*Kiss1r*gKO).
- *Kiss1r*gKO mice of both sexes are severely hypogonadal.
- *Kiss1r*gKO females, but not males, are obese with impaired glucose tolerance.
- *Kiss1r*gKO females show decreased VO_2 and VCO_2 , likely underlying their obesity.
- No sex differences were identified in *Kiss1* or *Kiss1r* expression in peripheral tissues.
- These data further highlight a key role for Kiss1r signaling in BW and metabolism.

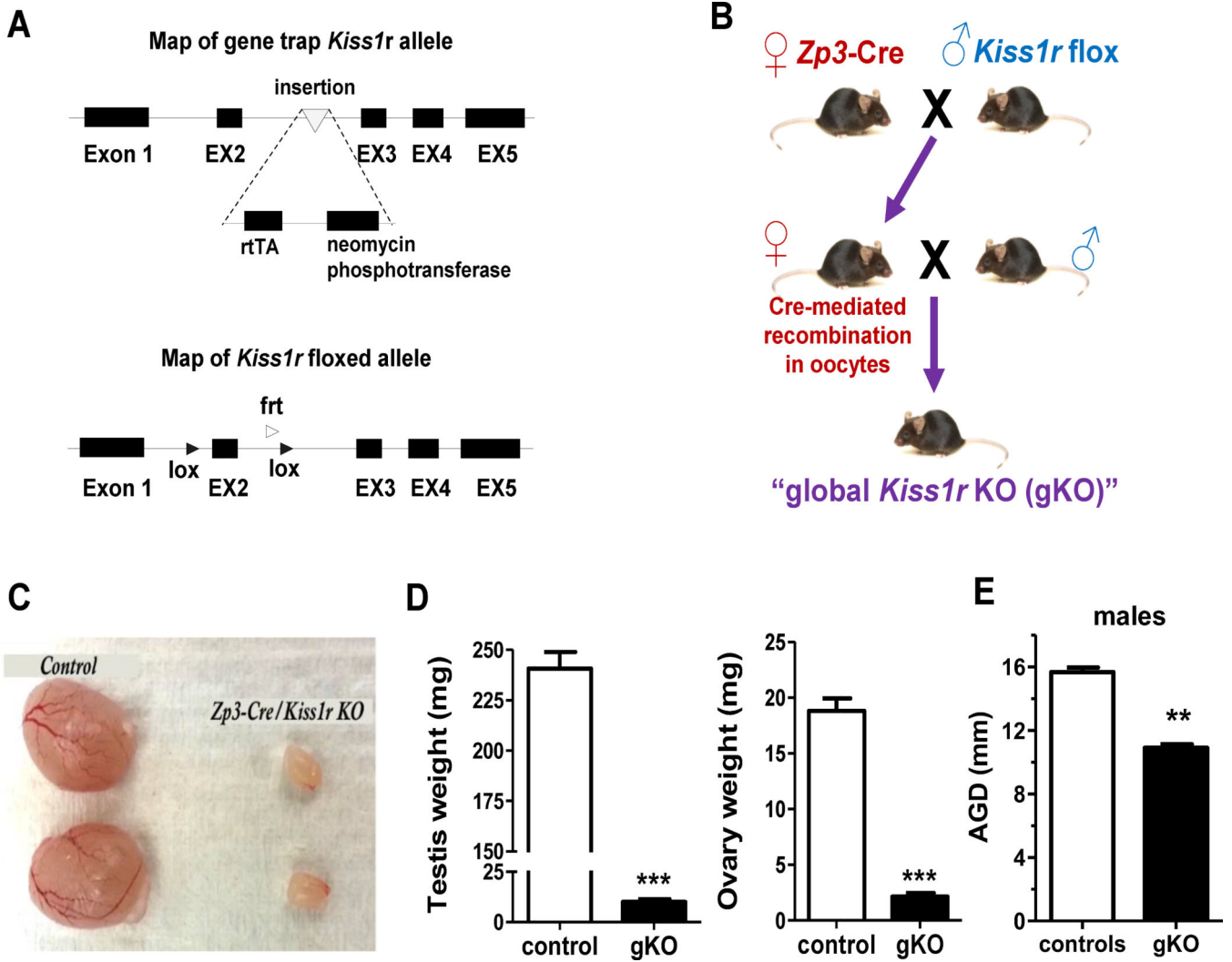


Figure 1: Development and reproductive characterization of a new global *Kiss1r* KO mouse made using Cre/lox technology. **A)** Map of the *Kiss1r* allele in the original gene-trap *Kiss1r* KO mouse line and the *Kiss1r* floxed line used in this study. **B)** Schematic of the simplified strategy for creating the new global *Kiss1r* KO (“gKO”) by crossing *Zp3-Cre* and *Kiss1r* floxed strains. **C)** Image of testes of adult gKO and WT males. **D)** Ovary and testes weights of adult gKO and control mice (n=6–9/genotype for each sex). **E)** AGD of adult male gKOs (n=10–15/genotype). **, p<0.01, ***p<0.001.

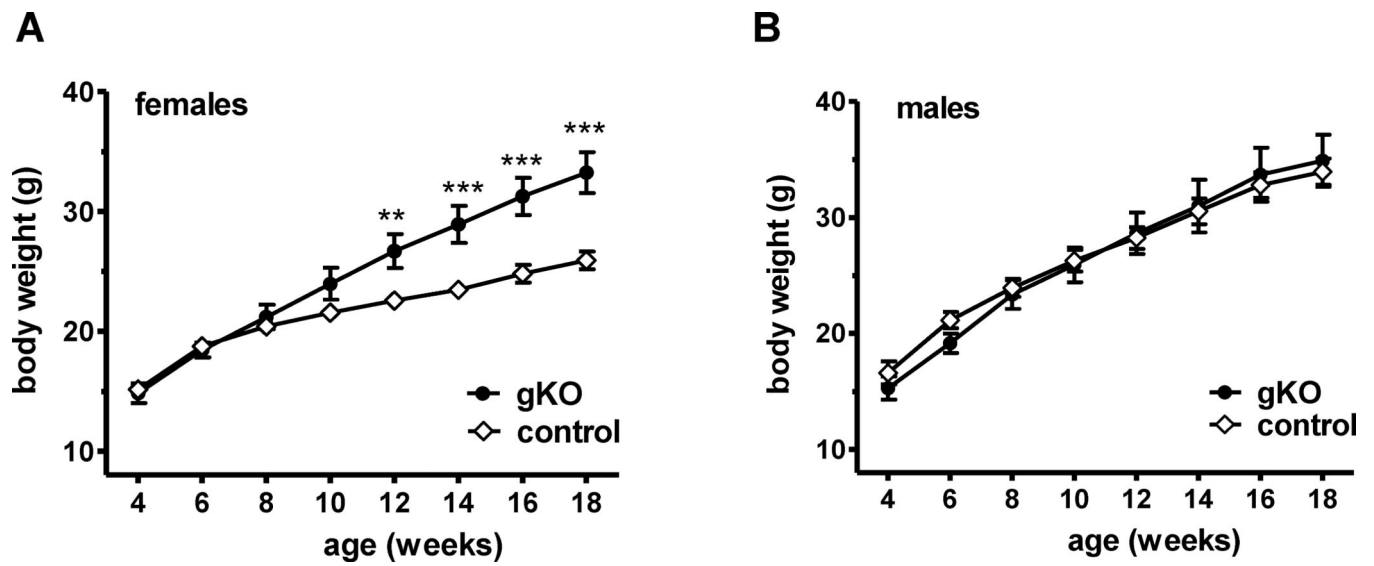


Figure 2: Body weights of female and male *Kiss1r* gKO mice made using Cre/lox technology. **A)** BWs of gKO and control female mice (n=9–15/genotype). **B)** BWs of gKO and control male mice (n=7–9/genotype). **, p<0.01, ***p<0.001.

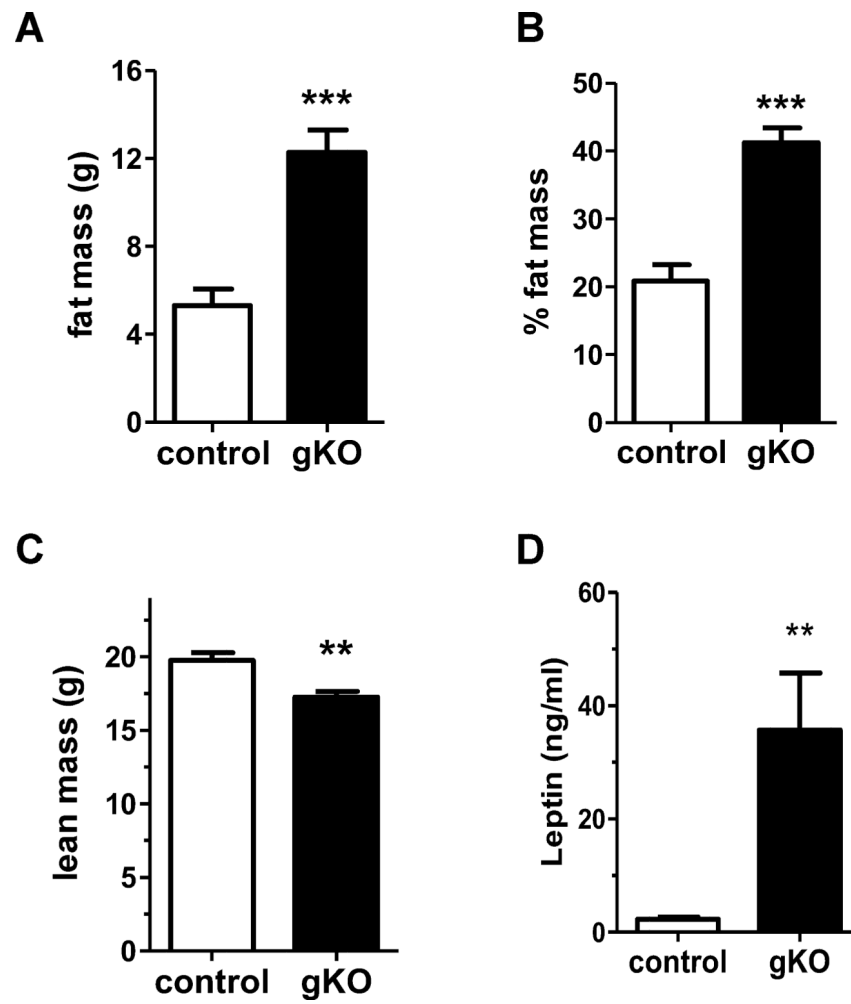


Figure 3: Body composition of adult female gKO and control mice. **A)** Fat mass, **B)** percent fat mass, **C)** lean mass, and **D)** blood serum leptin levels are all significantly altered in gKO females. n=6–10/genotype. **, p<0.01, ***p<0.001.

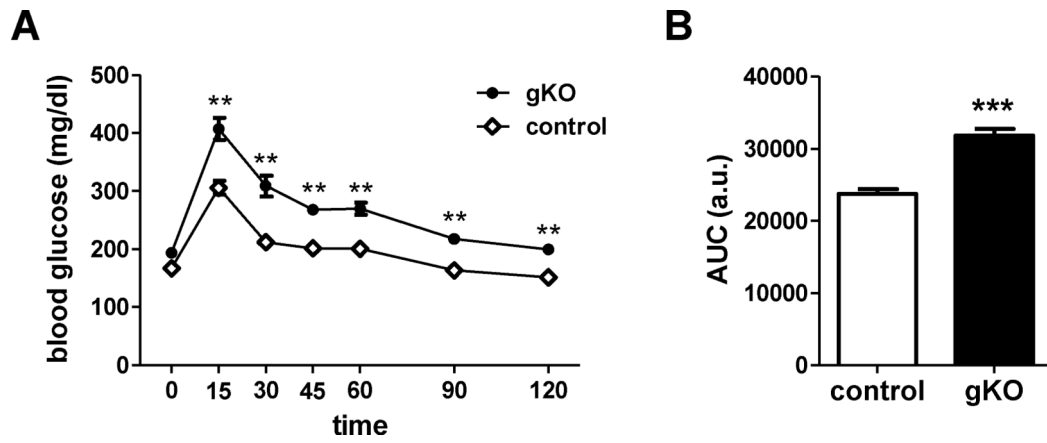


Figure 4:

A) Glucose regulation of adult gKO female mice is impaired in an ip glucose tolerance test (GTT). **B)** Area under the curve (AUC) for gKO and control mice during the GTT. n=10–12/genotype. **, p<0.01, ***p<0.001.

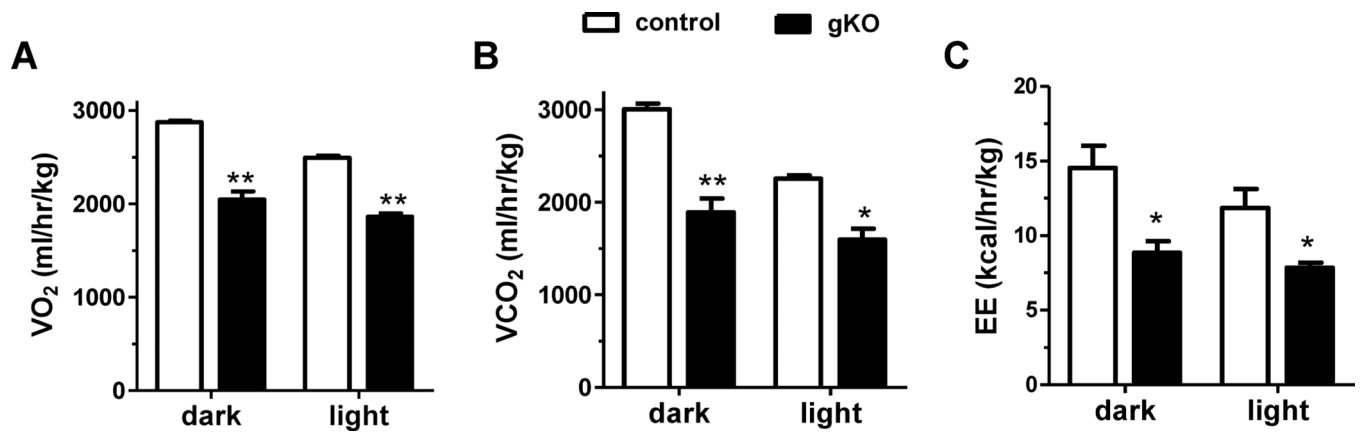


Figure 5: Metabolic rates and energy expenditure in controls versus gKO females. **A)** Oxygen consumption (VO₂), **B)** carbon dioxide production (VCO₂), and **C)** energy expenditure are all significantly decreased in gKOs in both the light and dark periods of the daily light cycle (n=3/genotype). *, p<0.05, **, p<0.01.

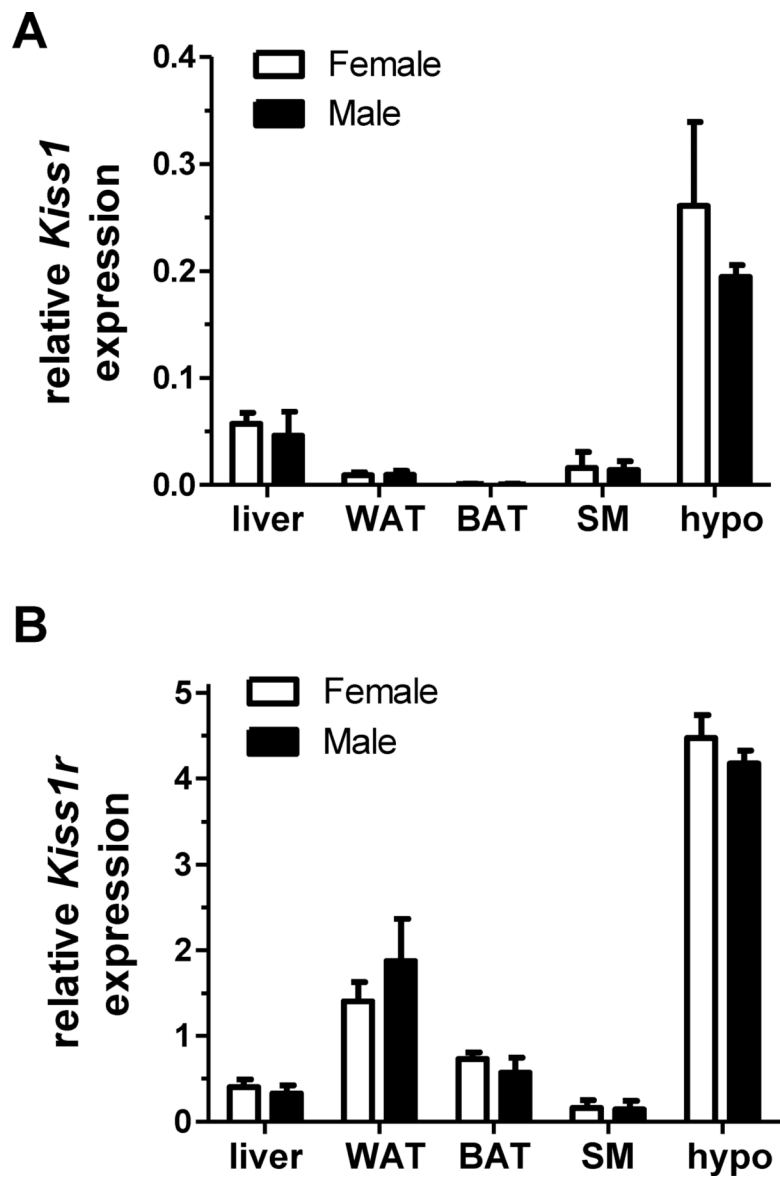


Figure 6: qPCR gene expression analysis of *Kiss1* and *Kiss1r* levels in several metabolic-relevant tissues, including brain and peripheral tissues, of normal adult GDX male and female mice. **A)** *Kiss1* levels in males and females. **B)** *Kiss1r* levels in males and females. There were no sex differences detected for either gene in any tissue examined. WAT, white adipose tissue; BAT, brown adipose tissue; SM, skeletal muscle; hypo, hypothalamus.