Deletion of Six3 in post-proliferative neurons produces weakened SCN circadian output, improved metabolic function, and dwarfism in male mice

ABSTRACT

Objective: The increasing prevalence of obesity makes it important to increase the understanding of the maturation and function of the neuronal integrators and regulators of metabolic function.

Methods: Behavioral, molecular, and physiological analyses of transgenic mice with Sine oculis 3 (Six3) deleted in mature neurons using the SynapsinIIe allele.

Results: Conditional deletion of the homeodomain transcription factor Six3 in mature neurons causes dwarfism and weakens circadian wheel-running activity rhythms but increases general activity at night, and improves metabolic function, without impacting pubertal onset or fertility in males. The reduced growth in 6-week-old Six3\textsuperscript{fl/fl-SynapsinIIe} (Six3\textsuperscript{fl/fl}) males correlates with increased somatostatin (SS) expression in the hypothalamus and reduced growth hormone (GH) in the pituitary. In contrast, 12-week-old Six3\textsuperscript{fl/fl} males have increased GH release, despite an increased number of the inhibitory SS neurons in the periventricular nucleus. GH is important in glucose metabolism, muscle function, and bone health. Interestingly, Six3\textsuperscript{fl/fl} males have improved glucose tolerance at 7, 12, and 18 weeks of age, which, in adulthood, is associated with increased % lean mass and increased metabolic rates. Further, 12-week-old Six3\textsuperscript{fl/fl} males have reduced bone mineralization and a lower bone mineral density, indicating that reduced GH levels during early life cause a long-term reduction in bone mineralization.

Conclusion: Our study points to the novel role of Six3 in post-proliferative neurons to regulate metabolic function through SS neuron control of GH release.

1. INTRODUCTION

Obesity is an increasing problem worldwide, with the World Health Organization estimating 1.9 billion adults were overweight in 2016, and 39 million children under the age of 5 were overweight or obese in 2020 [1]. Overweight and obesity are associated with increased risks of type 2 diabetes [2] and cardiovascular problems [3], among many other negative health outcomes. Although obesity is reversible and preventable through sustained changes in lifestyle, such approaches often fail. It, therefore, remains a priority to identify genes and pharmacological targets that regulate food intake and metabolic function, which are potential novel treatment strategies to help reduce the current worldwide epidemic of obesity.

Food intake and energy expenditure are regulated by the hypothalamus [4–6]. To maintain metabolic homeostasis, peripheral signals are integrated by hypothalamic nuclei, including the arcuate nucleus, periventricular nucleus (PeVN), and paraventricular nucleus (PVN). The function of these brain structures is aligned to the time-of-day and metabolic cues, such as meal hours. Time-of-day information is transmitted to the arcuate nucleus, PeVN, and PVN through direct projections from the suprachiasmatic nucleus (SCN). The SCN receives light information from the eye and translates it into neuronal and...
endocrine signals, providing day-length and light information to non-SCN brain areas and peripheral tissues. Disruption of SCN circadian time keeping or molecular clock function can negatively affect metabolic function and is associated with an increased risk of metabolic disease and diabetes [7–11]. Metabolic dysregulation can be induced through genetic approaches such as knock-out (KO) of molecular clock genes, including Bmal1, expression of the dominant-negative ClockΔ19 mutant [5,8], or light-induced circadian disruption. These relationships between metabolic function and circadian rhythms are further evidenced in studies focusing on feeding time, where time-restricted feeding, or eating earlier in the day, reduces weight gain and improves metabolic function [12]. This points to an interesting interaction between the circadian system, feeding, and metabolic status.

In addition to mis-timed light and poor eating patterns as risk factors in the development of overweight and obesity, genetic factors also contribute to the deregulation of metabolic function. Mutations in genes encoding neuropeptides, hormones, and their receptors that regulate hypothalamic function, including the genes encoding for POMC (pro-opiomelanocortin), leptin, and leptin receptor, cause obesity in rodents and humans [4]. A second set of genes pointing to the key role of dysregulated hypothalamic function as a cause of overweight and obesity are mutations in genes coding for transcription factors required for hypothalamic development and function, such as SIM1 [13,14]. A novel candidate gene involved in hypothalamic function is Sine oculis 3 (Six3) [15]. SIX3 is a homeodomain transcription factor required for retinal, forebrain, and pituitary development [16–24]. After embryonic development, Six3 retains a broad expression pattern in the brain [25], where it regulates radial glia maturation into ependymal cells [26], medium spiny neuron plasticity and striatal function [27–29], hypothalamic kisspeptin neuron function [30], as well as SCN function [25]. However, the potential role of Six3 in metabolism remains unexplored.

Based on the high expression of Six3 in the developing and adult male hypothalamus, and our previous work showing dysregulated hypothalamic function in female mice with Six3 deleted in post-proliferative neurons using a SynapsinIIα allele (Six3syn mice), we hypothesize that Six3syn males also have dysregulated hypothalamic function and impaired SCN circadian time keeping, which we predict will negatively impact glucose metabolism, increasing fat accumulation, and body weight.

2. MATERIALS AND METHODS

2.1. Mouse breeding

All animal procedures were performed according to the protocols approved by the Institutional Animal Care and Use Committees of University of California San Diego and Michigan State University, and were conducted in accordance with the Guide for the Care and Use of Laboratory Animals (National Research Council, 2011). Mice were maintained on a light/dark cycle of LD [12 h light, 12 h dark; lights on 6 AM (Zeitgeber ZT 0) to 6 PM (ZT12)], with ad libitum access to water and chow (Teklad S-2335 Mouse Breeder Irradiated Diet, Envigo). Animals were housed in polypropylene cages (25.4 x 20.0 x 12.7 cm) and caged in wire-mesh Pens with sawdust bedding. Male mice were housed with a virgin female (e−/−, e+/−, e+/+). All cage mates were heterozygous for their respective Cre−/−alleles. Six3 mutant mice were bred in-house to generate Six3tm2Gco, MGI 3693321 mice [31] were crossed with either B6.Cg-Tg (Syn1-cre)671Jxm/J, JAX #003966 mice. All females [34,35]. The number of pups per litter was recorded for the first 3 litters, as described previously [34].

2.3. Litter size assessment

For litter size assessment, virgin 8- to 12-week-old Six3tn/m and Six3t/m males were housed with Six3tn/m females [34,35]. The number of pups per litter was recorded for the first 3 litters, as described previously [34].

2.4. Quantitative real-time PCR

For RT-qPCR, the hypothalamus, pituitary, and liver were collected from males that were euthanized by CO2 overdose between 10AM (ZT4, 4 h after lights ON) and 1 PM (ZT7), except where otherwise stated. Tissues were snap frozen on dry ice and stored at −80 °C until RNA extraction. Total RNA was extracted using TRizol (Invitrogen). Genomic DNA was eliminated using the DNA-free kit (Applied Biosystems). CDNA was obtained by reverse transcription of total RNA using an iScript cDNA synthesis kit (Bio-Rad Laboratories). cDNA products were detected using an iQ SYBR Green Supermix (Bio-Rad Laboratories) on a qRT-PCR CFX real-time detection system (Bio-Rad Laboratories). qRT-PCR primers (Supplementary Table 1) were previously published [35–37]. Data were expressed as fold change using the 2−ΔΔCT method by normalizing Gnrh1 to Gapdh [38]. Data represent mean fold change ± SEM from a minimum of three mice for each data point.

2.5. Wheel-running behavior

At age 8–12 weeks, males were single housed in cages containing running wheels with magnets, and wheel revolutions were monitored using magnetic sensors. All cages were contained in a light-tight cabinet with programmable lighting and monitored for temperature and humidity. Food and water were available ad libitum during the entire experiment. Males were housed in polypolyethylene cages (17.8 x 25.4 x 15.2 cm) containing a metal running wheel (11.4 cm diameter), and locomotor activity rhythms were monitored with a VitalView data collection system (Version 4.2, MiniMitter, Bend OR) that integrated in 6 min bins the number of magnetic switch closures triggered by half wheel rotations. Running wheel activity was monitored for 4 weeks on LD (light12h:dark12h), where week 1 was considered as adaptation to a novel environment, and the two weeks prior to constant darkness (DD), were used as control baseline activity in LD. Subsequently, mice were monitored for 4 weeks in DD, with wheel running data analyzed from weeks 2 and 3 in DD. Cage changes were scheduled at 3-week intervals. Wheel-running activity was analyzed using ClockLab Analysis (ActiMetrics) by an experimenter blind to experimental group. Circadian period was estimated by constructing a ClockLab-generated least-squares regression line through a minimum of 13 daily activity onsets. Daily onset and offset of
activities, defined as a period of 5 h of activity following 5 h of inactivity (onset) or a period of 5 h of inactivity following 5 h of activity (offset), were used to calculate the duration of the active phase (alpha). Chi² periodograms [39] were generated for periods from 0 to 36 h, with a significance criterion set at 0.001. Any mice that did not exhibit a significant peak between 18 and 36 h were deemed arrhythmic and were not included in analyses, and the maximum amplitude was reported as Qp. Activity profiles were generated for weeks 2–4 in DD using the circadian period (tau) estimated from the chi² periodogram for the same time interval. Total daily counts were calculated over 24 h, during habituation, LD, and DD.

2.6. Ex vivo tissue recordings of Per2::LUC expression

For SCN, pituitary, epididymis, and liver explant studies, male Per2::LUC circadian reporter mice were used [40]. Males were sacrificed at ZT3–4 via isoﬂurane inhalation and cervical dislocation. The pituitary, liver, and brain were removed immediately and placed in ice-cold HBSS for approximately 30–60 min. Using a Vibratome (Leica), coronal brain sections of 300 μm were collected, and the SCN was dissected from the slices in ~2 × 2 mm squares. An SCN, ~3 × 2 mm liver section, one epididymis, or one pituitary was placed individually on a 30 mm MultiCell membrane (Millipore-Sigma) in a 35 mm cell culture dish containing 1 mL Neurobasal-A Medium (Gibco) with 1% Glutamax (Gibco), B27 supplement (2%; 12,349–015, Gibco), and 1 mM luciferin (BD Biosciences). The lid was sealed to the plate using vacuum grease to ensure an air-tight seal. Plated tissues were loaded into a LumiCycle luminometer (Actimetrics) inside a 35 °C non- humidified incubator at ZT6–6.5, and recordings were started. The bioluminescence was counted for 70 s every 10 min for 6 days (day 1–day 7 of recording time). Per2:LUC rhythmic data were analyzed using LumiCycle Analysis software (Actimetrics) by an experimenter blind to the experimental group. Data were detrended by subtraction of the 24 h running average, smoothed with a 2 h running average, and fitted to a damped sine wave (LM Fit, damped). Period was defined as the time in hours between the peaks of the fitted curve. Amplitude was defined as the amplitude of the fitted sine wave [41].

2.7. Dual energy X-ray absorptiometry (DEXA)

Body composition was determined in 12-week-old Six3<sup>/fl</sup> and Six3<sup>/yn</sup> males by DEXA. Mice were fasted for 4–6 h and then anesthetized (ketamine 100 mg/kg, xylazine 10 mg/kg). Body weight (BW) was measured, and lean muscle mass and fat mass were determined by scanning with a GE Lunar Pixi Densitometer Machine (n = 6 group).

2.8. Weekly growth analyses

Group housed males were weighed once weekly from 3 to 12 weeks of age. Length measurements were done on isoflurane anesthetized males using a caliper. The measurements were taken from the tip of the nose to the base of the tail in 3, 4, 6, 8, 12-, and 16-week-old males.

2.9. Hormone assays and glucose tolerance tests (GTT)

GTTs were performed in adult Six3<sup>/fl</sup> and Six3<sup>/yn</sup> male mice on a standard chow diet at 7-, 12-, and 18-weeks of age. Mice were fasted for 6 h before GTT with free access to water. Blood glucose was then measured using a handheld glucometer (One Touch UltraMini, LifeScan, Inc.) just before IP glucose injection (time 0; 2 g/kg BW in saline) and subsequently at 15, 30, 45, 60, 90, and 120 min post-administration using 1 g/kg BW glucose in saline. For all other serum hormone analyses, mice were euthanized by isoflurane overdose between ZT4 to ZT7 and blood collected from the abdominal aorta. Blood was allowed to clot for 1 h at RT, then centrifuged (RT, 15 min, 2600 g), and serum stored at –20 °C before Luminex analysis for GH, LH, and FSH on MILLIPLEX MAP Mouse Pituitary Magnetic Bead Panel (Millipore Sigma #MIP7MA-T49k). Coefficients of variance (CVs) were based on the variance of samples in the standard curve run in duplicate or triplicate. Lower detection limits: LH = 5.6 pg/mL, CV < 15%; FSH = 25.3 pg/mL, CV < 15% GH = 4.88 pg/mL, GH < 10%. Samples were run in singlets.

2.10. Metabolic and locomotor analyses

Indirect calorimetry was performed on 18- to 21-week-old Six3<sup>/yn</sup> males on a standard chow diet using a 12-cage equal flow CLAMS calorimeter system (Columbus instruments) coupled with photo-sensors to detect movement. Each cage has internal dimensions of: width (4 cm) x depth (20 cm) x height (13 cm). Males were habituated to the metabolic cages (single-housed) for 2 days prior to data acquisition (n = 6/genotype). O<sub>2</sub> consumption (VO2) and CO<sub>2</sub> production (VCO2) were measured every 12 min in each cage. Respiratory exchange ratio (RER) was calculated as the quotient of VCO2/VO2. Locomotor activity was measured in 1 min intervals by photo-sensors, with the bottom row measuring horizontal movement (total = total number of photobeams broken; ambulatory = number of consecutive beams broken to exclude non-ambulatory movements like grooming) and the upper row measuring vertical movement (includes rearing, reaching the drinking tube, walking on top of the food hopper). In addition, feeding was measured in 12 min intervals and calculated for hourly feeding. Cage tops were equipped with infrared sensors (TSE InfraMot; TSE Systems) to detect body-heat image and its spatial (3 dimensional) displacement over time to assess general locomotor activity, measured as arbitrary ‘InfraMote- units’. Owing to high fluctuations of the respiratory rate readings, CO<sub>2</sub> and O<sub>2</sub> records from each mouse were subjected to the filtering process. The median and standard error were calculated for every 2.5-hour interval (i.e., 12 observations), and the observations within the range of median ± standard error were selected and used for further analysis. There was a total of 306 raw observations of O<sub>2</sub> and CO<sub>2</sub> per cage (mouse), and on average 210 observations were remaining after the filtering process. To fit the sinusoidal curve, each filtered dataset was then fitted with equation Y = A sin (B t + C) + D by using optimization functions embedded in the SciPy package in Python 3.6 (Python Software Foundation). Model parameter A indicates the amplitude, B indicates the frequency, C indicates the phase, and D indicates the mean. From the frequency and phase, the time for the VO2 or VO2 to reach maximum (T<sub>max</sub>) was calculated. T<sub>max</sub> was adjusted to have a range between 12 and 36 h after the onset of the start of data collection. The average R² values of the curves were 0.43 and 0.45 for O<sub>2</sub> and CO<sub>2</sub> datasets, respectively. The parameters derived from curve fitting were analyzed by PROC TTEST in SAS 9.4 (SAS Institute Inc., USA). The model included the parameter (amplitude, mean, or T<sub>max</sub>) as a variable and the mouse type (i.e., Six3<sup>/fl</sup> and Six3<sup>/yn</sup>) as a class.
and 1x penicillin-streptomycin (Life Technologies/Invitrogen) in a DMEM (Mediatech), containing 10% fetal bovine serum (Gemini Bio), NIH3T3 cells (American Type Culture Collection) were cultured in AVP and VIP staining in SCN).

DNA transfected into cells, we systematically equalized plasmid used was mouse Six3/pcDNA overexpression plasmid (200 ng/well, values were normalized to \[b\] were harvested 24 h after transfection in lysis buffer \[100 mM potassium phosphate (pH 7.8) and 0.2% Triton X-100\], Luciferase values were normalized to \[b\] -galactosidase to control for analysis by Tukey or Bonferroni as indicated in figure legends, with \( p < 0.05 \) to indicate significance. First peak phase relationships of Per2:LUC timing were analyzed in R via a Circular Analysis of Variance High Concentration F-Test, with a corrected confidence level of \( p < 0.01667 \) to account for family-wise error. Wheel-running activity was analyzed via a two-way repeated-measures ANOVA.

### 3. RESULTS

#### 3.1. Six3<sup>39m</sup> males have impaired SCN function, but intact fertility

We recently showed that the Prosapsin<sup>39m</sup>-allele recombines the Six3<sup>39m</sup>-allele throughout the brain including the SCN, impairing SCN circadian output in Six3<sup>39m</sup> females [25]. To determine whether circadian function was also weakened in Six3<sup>39m</sup> males, we placed Six3<sup>39m</sup> and Six3<sup>39m</sup> males on running-wheels (Figure 1A). To limit the number of mice in the male wheel-running study, as it was an extension of our previous study in Six3<sup>39m</sup> females, we decided to estimate the number of mice required to reach the lower power of 60% based on chi square amplitude (Op) values of our previous publication studying wheel-running of Six3<sup>39m</sup> females [25]. Our ad-hoc power analysis indicated that a total of 16 mice would allow us to achieve more than 60% of power. We first evaluated Six3<sup>39m</sup> male activity on the running-wheels, as assessed through average wheel revolutions. Six3<sup>39m</sup> males had comparable activity to Six3<sup>39m</sup> during the lights ON and OFF of the habituation period (Hab. ON and Hab. OFF, Figure 1B), and during the baseline (LD) activity period following the habituation period (LD ON and LD OFF, Figure 1B). Running wheel activity is a known rewarding activity in rodents [45,46]. To determine whether the non-significant trend toward reduced activity of Six3<sup>39m</sup> males during lights OFF (Figure 1B) reflected a global reduction in reward function, we performed a sucrose preference test in a new cohort of mice. Six3<sup>39m</sup> mice presented with a sucrose preference over water comparable to Six3<sup>39m</sup> mice, indicating overall normal reward processing of Six3<sup>39m</sup> males (Figure 1C, 3-way ANOVA, effect genotype F (1, 50) = 1.646, \( p = 0.206 \). To determine whether the endogenous free-running period of the SCN was changed in Six3<sup>39m</sup> mice, we evaluated the wheel-activity rhythm period. Six3<sup>39m</sup> mice had a lengthened free-running period in DD as compared to Six3<sup>39m</sup> (Tau, Figure 1D, 2-way ANOVA, effect of genotype F (1, 22) = 14.51, \( p < 0.001 \). It should be noted that we observed a surprisingly large variation in Tau in the controls (Figure 1D), a variation that was even more pronounced in the Six3<sup>39m</sup>. Indeed, as can be observed on the individual actigraphy data from 6 controls and 8 Six3<sup>39m</sup> males (Supplementary Figure 1), the activity onset of the controls during DD was relatively variable, suggesting the Six3<sup>6th</sup> allele by itself impacts SCN function. Interestingly, as we had previously observed for Six3<sup>39m</sup> females [25], 6/8 Six3<sup>39m</sup> males on DD had very variable activity periods, where they initially trended to have a period shorter than 24 h, but after variable times on DD, suddenly switched to a longer Tau. Owing to the low activity profile of some of the Six3<sup>39m</sup> males (Supplementary Figure 1, Six3<sup>39m</sup>), we were unable to analyze the data from some of the mice. The great variation in wheel-activity rhythm and activity levels of the Six3<sup>39m</sup> males led to a great variation in the shortening/lengthening in Tau in DD of Six3<sup>39m</sup> males, a
variation that also resulted in variable activity phase (alpha), which was comparable in Six3fl/fl and Six3syn mice (Figure 1E, 2-way ANOVA, effect of genotype F (1,12) = 0.01, p < 0.92). To determine whether the lengthened Tau in Six3syn was associated with weakened circadian timekeeping in the SCN, we evaluated circadian rhythm strength by chi square. Six3syn had reduced chi square amplitude (Qp) in LD (Figure 1F) and DD (Figure 1F,G). SCN circadian output precision and strength are regulated by numerous factors, including SCN neuron connectivity. To determine whether the change in SCN output might be through an effect of SIX3 on the expression of SCN peptides, we evaluated SCN expression of somatostatin (SS), arginine vasopressin (AVP) and vasoactive intestinal polypeptide (VIP) by immunohistochemistry. No significant difference was observed in the number of SCN neurons expressing SS (Figure 1H,K), AVP (Figure 1I,L) or VIP (Figure 1J,M) in the Six3syn males as compared to controls.
To determine whether the reduced SCN circadian precision was associated with changes in the molecular clock within SCN neurons, we analyzed the capacity of SIX3 to modulate the molecular clock genes Per2 and Bmal1 through in vitro transient transfection studies. We recently showed that SIX3 overexpression in vitro increased Per2-luciferase expression through ATTA-sites in the Per2 regulatory region [25]. To determine whether SIX3 also regulates Bmal1 expression, we analyzed the Bmal1 regulatory region for potential SIX3 binding sites (ATTA-like sites) and identified five potential SIX3 binding sites (Supplementary Table 2). One potential binding site, located at −537 bp, was not present in either plasmid creator’s original sequence or in any build of the mouse genome. As such, we modified the plasmid back to its correct sequence, leaving four potential homeodomain binding sites. To assay for SIX3 regulation of Bmal1 expression, we transiently transfected NIH3T3 cells with the regulatory region of Bmal1 driving luciferase (Bmal1-luciferase), with SIX3 or empty vector (pcDNA). SIX3 overexpression increased Bmal1-luciferase expression 2.3-fold (Figure 2A). To determine whether the enhanced expression of Bmal1-luciferase was mediated through the putative SIX3 binding sites, we created four plasmids with single ATTA-like sites mutated, in addition to one plasmid with all four ATTA-like sites mutated (ΔATTA). All the mutated ATTA sites impacted SIX3 regulation of Bmal1-luciferase expression (Figure 2A).

It is well established that changes in Bmal1 and PER2 impact circadian timekeeping. To determine if the capacity of SIX3 to regulate core clock gene expression in vitro translated into changes in circadian timekeeping function in SCN tissue, we generated triple transgenic mice, crossing Six3syn::Per2::LUC mice with Per2:LUC knock-in reporter mice ([25]). To determine whether SIX3 also regulates Per2:LUC circadian period in explants of SCN and epididymis. A) Transient transfections of NIH3T3 cells with a reporter plasmid containing the mouse Bmal1 regulatory region driving the expression of firefly luciferase (Bmal1-luciferase) with SIX3 or empty vector (pcDNA). SIX3 overexpression vector (200 ng, green) or pcDNA 200 ng, white). Location of site-directed mutagenesis of ATTA and ATTA-like sites in the mouse Bmal1–luciferase plasmid are indicated with the Δ. For ΔATTA, all ATTA sites in the regulatory region have been mutated (see Supplementary Table 2). Statistical analysis by one-way ANOVA, with Dunnett’s post-hoc analysis as compared to Six3/−/Bmal1−/− luciferase, *, p < 0.05; **, p < 0.01; ***, p < 0.001, n = 5–6 in triplicate. B) Example recording of SCN Per2:LUC circadian period. Histogram of Per2:LUC period and amplitude in the C) SCN, D) pituitary, E) liver and F) epididymis from Six3fl/fl::Per2::LUC and Six3myr::Per2::LUC males. Statistical analysis by Student’s t-test, *, p < 0.05; **, p < 0.01, n = 3–6.

Figure 2: SIX3 regulates Bmal1-luciferase expression in NIH3T3 cells and Per2:LUC circadian period in explants of SCN and epididymis. A) Transient transfections of NIH3T3 cells with a reporter plasmid containing the mouse Bmal1 regulatory region driving the expression of firefly luciferase (Bmal1-luciferase) with SIX3 or empty vector (pcDNA 200 ng, green). B) Example recording of SCN Per2:LUC circadian period. Histogram of Per2:LUC period and amplitude in the C) SCN, D) pituitary, E) liver and F) epididymis from Six3fl/fl::Per2::LUC and Six3myr::Per2::LUC males. Statistical analysis by Student’s t-test, *, p < 0.05; **, p < 0.01, n = 3–6. www.molecularmetabolism.com
Six3<sup>3yn</sup> males have increased % lean mass, improved GTT, and increased metabolic rate

To further explore the ~50% reduction in body weight of adult Six3<sup>3yn</sup> males compared to Six3<sup>fl/fl</sup> males (Figure 3B), we used dual energy X-ray absorptiometry (DEXA) to measure the body composition of 12-week-old male mice. As compared to Six3<sup>fl/fl</sup> males, Six3<sup>3yn</sup> males had significantly less total fat mass (Figure 4A), a lower % fat mass (Figure 4B), and lean less mass (Figure 4C), but increased lean mass as a percentage of total body mass (Figure 4D). To determine whether the increase in % lean mass was associated with improved glucose tolerance, we performed a glucose tolerance test (GTT) at 7, 12, and 18 weeks of age. No significant difference in fasted basal blood glucose levels were identified (Figure 4E). In contrast, at all the ages studied, Six3<sup>3yn</sup> males showed reduced blood glucose levels compared to Six3<sup>fl/fl</sup> (Two-way ANOVA, repeated measures). Effect of genotype: 7-weeks-of-age F (1, 126) = 56.47, p < 0.0001; 12-weeks-of-age F (1, 42) = 58.20, p < 0.001; 18-weeks-of-age F (1, 42) = 86.62, p < 0.0001.

We next examined whether hypophagia contributed to the reduced fat mass in these mice by using the comprehensive Lab Animal Monitoring System (CLAMS) metabolic cages. Consistent with a higher metabolic demand for a smaller animal [50], Six3<sup>3yn</sup> males had a higher metabolic rate than Six3<sup>fl/fl</sup> (Figure 5A-E), and ate more chow (Figure 5F,G), without eating for longer time periods (Figure 5H), but had normal energy expenditure (Figure 5I). In addition, Six3<sup>3yn</sup> males had increased locomotor activity, as evaluated using beam breaks in the metabolic cages (Figure 5J,K), and repetitive behavior during the dark phase compared to Six3<sup>fl/fl</sup> (Figure 5L). Consistent with the increased nocturnal activity (Figure 5J,K), K: Two-way ANOVA interaction genotype with time [F (22, 220) = 2.278, p = 0.014, n = 6], Six3<sup>3yn</sup> males had increased VO2 and VCO2 at night as compared to Six3<sup>fl/fl</sup> males (Figure 5A,D). Changes in locomotor activity and weakened SCN output can change the timing of peak VO2 and VCO2 rates [51]. To determine the time of day of peak VO2 and VCO2, we smoothed the VO2 and VCO2 data, and used a sinusoidal model to identify the rhythm phase (Figure 5M, N, example V2O2 data illustrated, see Supplementary Figs. 2–5 for individually smoothed data sets and model fitting). We found no change in the phase (peak time) of VO2 (Figure 5O) or VCO2 (Figure 5P). To determine whether the dysregulated metabolic function might be driven, in part, by Synapsin<sup>3yn</sup> recombination of the Six3<sup>ox</sup> allele in peripheral tissues, we tested for Synapsin<sup>3yn</sup>-driven Six3<sup>3yn</sup>-allele recombination. We confirmed that in males Synapsin<sup>3yn</sup> targets the hypothalamus and pituitary (Figure 5Q) and also identified recombination in the tests. No recombination was seen in white and brown adipose tissue, liver, tail or the pancreas (Figure 5O). Of these tissues, Six3 is known to be expressed in the hypothalamus, cortex, and pituitary (Figure 5Q, *) [18].

3.3. Six3<sup>3yn</sup> males have increased SS in the periventricular nucleus (PeVN)

Within the brain, metabolic control is regulated by the hypothalamus, where genes coding for neuropeptides known to be involved in metabolic homeostasis include Npy (neuropeptide Y), Kiss1 (kisspeptin), Ghrh (growth hormone releasing hormone), and Sst (somatostatin) [4]. Npy, Kiss1, and Ghrh-expressing neurons are primarily located in the arcuate nucleus, whereas SS neurons are located in the PVN and PeVN (Figure 6A–C). The arcuate nucleus, PVN, and PeVN are targeted by the Synapsin<sup>3yn</sup>-allele [25], and express Six3<sup>3yn</sup> (Figure 6A–C). To determine what neuronal population(s) might be involved in the altered metabolic parameters of the Six3<sup>3yn</sup> males, we performed qPCR on the entire hypothalamus. At 6 weeks of age, but not at 12 weeks of age, Six3<sup>3yn</sup> males had a significant increase in hypothalamic expression of Ss and Npy, but no significant change in Tshb, Ghrh (Figure 6D–G), or receptors integrating metabolic status were observed (Figure 6H).

The limitation of using whole hypothalamus extracts for qPCR is the incapacity to detect moderate or small gene expression changes in discrete neuronal populations. To separate out the neuronal
population(s) contributing to the reduced growth and increased metabolic rate in Six3syn males, we conditionally deleted Six3 from kisspeptin neurons (Six3kiss). We found that Six3kiss males had comparable body weight to control (Six3fl/fl) males from 3 to 12 weeks of age (Figure 6I), indicating the significant change in body weight developing after 4 weeks of age in Six3syn males is unlikely to be driven by Six3 in kisspeptin neurons. We therefore shifted our focus to AVP and SS expressing neurons of the PVN and PeVN which are known to regulate metabolic status and the growth axis [52,53]. Using immunohistochemistry, we found that Six3syn males and Six3fl/fl controls had a comparable number of SS-expressing neurons in the PVN (Figure 6J, M), but Six3syn males had a greater number of SS-expressing neurons in the PeVN compared to control mice (Figure 6K, N). The increased number of SS-expressing neurons in the PeVN provides a mechanism to regulate pituitary release of GH, a hormone critical in growth and metabolism. We also evaluated AVP expression in the PVN, as AVP neurons in the PVN have been associated with neuroendocrine regulation of metabolic function [53,54], and we previously found SIX3 to be a potent activator of Avp expression [25]. We did not, however, detect a change in AVP cell numbers in the PVN of Six3syn males (Figure 6L, O).

3.4. Six3syn males have a dysregulated growth axis causing reduced body size and bone demineralization
SS is a known repressor of GH release (Figure 7A). To determine if the increase in SS in Six3syn males impacted Gh mRNA, we performed qPCR of mRNA extracted from whole pituitary. By 12 weeks of age, Six3syn mice exhibited a significant decrease in expression of Gh (Figure 7B). In contrast, GH release was significantly increased at 12 weeks of age (Figure 7C). To further explore how the changes in body weight might be caused by abnormal GH release, we used qPCR to evaluate two liver transcripts regulated by GH: Igf1 (insulin like growth factor 1) and Ghr (growth hormone receptor). We did not find a significant change in either Igf1 (Figure 8A) or Ghr (Figure 8B) in the adult Six3syn liver. Despite these non-significant changes of Igf1 expression in the liver, it remains possible the IGF1 protein is reduced, as Six3syn bone mineral count (Figure 8C) and bone mineral density (Figure 8D), were significantly reduced (see Figure 7A for summary of the growth axis).

4. DISCUSSION
Here, we demonstrate the novel role of SIX3 in the brain: conditional KO of Six3 in post proliferative neurons (Six3syn) is associated with reduced growth, increased lean mass, improved glucose tolerance, and weakened circadian wheel-running rhythms. These studies show for the first time the involvement of SIX3 in neuronal control of the growth axis and metabolic function and identify Six3 as a gene that warrants further studies to focus on PeVN SS neurons and neuronal control of GH release.

4.1. Reduced growth and dwarfism in Six3syn males
The most striking phenotype of the Six3syn males was their significant reduction in growth starting at 4 weeks of age, which was associated with increased % lean mass and reduced fat mass. As we recently described, conditional KO of Six3 in kisspeptin neurons (Six3kiss) resulted in reduced fertility [30]; therefore, we first tested whether this metabolic deregulation was caused by loss of SIX3 in the arcuate nucleus. We did not find any significant changes in arcuate nucleus mRNA expression for peptides or receptors regulating or sensing

Figure 4: Six3syn males have increased % lean body mass and improved glucose tolerance test. DEXA measurements of Six3syn males fed standard chow had reduced A) total and B) % fat mass, C) reduced total lean mass, and D) increased % lean mass at 12 weeks of age. E) Glucose tolerance test (GTT) results (blood glucose after IP glucose challenge) in 7, 12, and 18-week-old males. Statistical analysis by (A–D) Student’s t-test and (E) two-way ANOVA, repeated measures. *p < 0.05, **p < 0.01, ***p < 0.001, n = 4–6.
metabolic status, such as leptin receptor, insulin receptor, or the orexigenic Pomc, whereas a small but significant increase in the orexigenic Npy (neuropeptide Y) was detected at 6, but not 12 weeks of age [52,55–58]. Sucrose preference test performance was unaltered in Six3syn males. Indeed, the hyperphagia of Six3syn males correlated with increased metabolic rate, as expected for smaller, hyperactive animals [50]. These findings indicate that the reduced growth was not caused by dysregulated arcuate nucleus function. Rather, our data point to a novel role of SIX3 in PeVN SS neurons. A balanced release of arcuate GHRH, which defines the GH amplitude [59] and PeVN SS, which drives troughs of GH release [60,61], are well known to be required for normal growth, and suppression of GH in prepubertal mice is a leading cause of dwarfism [62]. Ideally, we would have studied the pulsatile release of GH in the Six3syn males; however, as GH pulsatile release in males is characteristic of one GH pulse per 2–4 h [37], we decided not to collect serial blood samples from Six3syn males, as their significantly smaller size, and resulting smaller blood volume as compared to controls, made this approach unfeasible in this mouse line. Alternatively, we compared GH levels at a single time point to SS and GHRH expression in the brain as an indicator of how GH release was impacted in Six3syn males. In agreement with the known repressive role of SS on GH release (Figure 7A), we found that Six3syn males had increased expression of Ss mRNA in the hypothalamus at 6-weeks of age, an increase that trended in 12-week-old males on the transcriptional level (Ss, p = 0.076) and was significantly increased when counting SS expressing neurons in the PeVN. The significant increase in Ss in the 6-week-old Six3syn males was associated with a significant reduction in Gh transcript, whereas circulating levels of GH was comparable to controls. The comparable levels of GH between Six3fl/fl and Six3syn might be caused by the timing of blood collection, potentially happening during GH troughs. This possibility is supported by the reduction in Gh transcript, which is not pulsatile, and which correlated with statured growth in young adult males, and dwarfism in adulthood. Interestingly in 12-week-old Six3syn males GH levels were

Figure 5: Six3syn males have increased metabolic rates. CLAMS cage assessment of feeding, metabolism, and locomotor activity in 12-week-old Six3syn males fed standard chow. A, B) CO2 production (VCO2), C, D) O2 consumption (VO2), E) respiratory exchange ratio (REx), F–H) feeding, I) energy expenditure, J, K) locomotor activity, and L) stereotypic repeated behavior. M, N) Example of smoothing and sine curve fitting to the VO2 data. O) VO2 and P) VCO2 phase (time of day of first peak) as estimated using the sine curve fitting to the smoothed data. Q) PCR to assay SynapsinCre+ driven recombination of the Six3lox allele in males. * indicated tissues known to express Six3. Abbreviations: WAT: white adipose tissue, BAT: brown adipose tissue. Statistical analysis by two-way ANOVA, repeated measures (A, C, E-K), and (O, P) Student’s t-test. **p < 0.01, *** p < 0.001. n = 5–6.
higher than controls. This seemingly contradictory finding can be explained by two independent mechanisms. First, it is again possible the single blood collection time point led us to collect blood in Six3\textsuperscript{fl/fl} only during GH troughs, whereas half of the blood samples collected from the Six3\textsuperscript{syn} were randomly collected during GH peaks. A second more plausible explanation is that the trending reduction in liver Igf1 promoted GH secretion from the pituitary [63,64], counteracting the increased inhibitory tone by SS on somatotrophs. Although we did not measure circulating IGF1 in this study, Six3\textsuperscript{syn} liver Igf1 mRNA levels trended downward, and two functional parameters enhanced by IGF1 were significantly reduced (bone mineralization and muscle mass, Figure 7A). Thus, we can infer from our data that adult Six3\textsuperscript{syn} males likely had impaired IGF1 feedback to the brain and pituitary, leading to increased GH release in adulthood, despite increased SS in the PeVN. Taken together, these findings support the known differential role of GH in early life versus adult and aging rodents [65]. Increased GH release in adulthood can be driven by reduced IGF1 negative feedback to the hypothalamus and somatotrophs. The dysregulated GH release in Six3\textsuperscript{syn} would also be expected to change expression of additional liver genes. What such genes are and how they contribute to the dysregulated metabolic function would be of great interest to explore in the future. Indeed, ~90% of sexually dimorphic genes in the liver are regulated by pulsatile GH patterns [66]. Thus, additional studies exploring how GH release patterns differ in Six3\textsuperscript{syn} males and females and how they contribute to changes in liver function would be of interest but are outside the scope of the present work.

Another important role of GH is its direct regulation of adipocyte function through the GH receptor. GH binding to its receptor promotes breakdown of triglycerides, thus increasing energy availability, while concomitantly suppressing the uptake of circulating lipids into adipocytes. Consistent with the increased GH release in 12-week-old Six3\textsuperscript{syn} males was their significant reduction of fat mass and increased % lean mass. The increase in % lean mass, along with the known role of GH in regulating glucose metabolism, resulted in improved glucose tolerance.
in *Six3<sup>syn</sup>* males as compared to controls, an improvement that increased with age. Taken together, these data point to a novel role of SIX3 in regulating the growth axis through regulation of SS neurons. It will be of interest in future studies to specifically delete Six3 in PeVN neurons in prepubertal and adult males to better understand the differential role of SIX3 in juvenile and adult animals in the regulation of the growth axis, fat versus lean body mass, and metabolic function.

### 4.2. Weakened SCN output contrasts with increased locomotor activity and metabolic rate in *Six3<sup>syn</sup>* males

SIX3 is required for SCN development [24] and for SCN circadian timekeeping in females [25]. Consistent with these previous results in female mice, we found that *Six3<sup>syn</sup>* males had weakened SCN output as measured by wheel-running activity rhythms. Interestingly, although the initial circadian period in DD was not significantly lengthened in *Six3<sup>syn</sup>* as compared to *Six3<sup>fl/fl</sup>* mice, 6 out of 8 *Six3<sup>syn</sup>* males showed a lengthening of period from <24h to >24h during DD, and this period lengthening was reflected in PER2:LUC recordings of SCN explants. These data are consistent with our previous finding of a general lengthening of SCN circadian period in *Six3<sup>fl/fl</sup>* females, attributable to a direct action of SIX3 on the Per2 promoter [25]. We show here that SIX3 also promotes Bmal1-luciferase expression *in vitro* and identify 4 ATTA sites required for this transcriptional activity. An additional contributing factor to the lengthening in SCN period in the *Six3<sup>syn</sup>* is a possible change in neuropeptide expression. *In vitro*, SIX3 is an activator of the Avp-luciferase, but not the Vip-luciferase promoter, whereas it is unknown if SIX3 can directly regulate Ss expression.

Using immunohistochemistry, we did not observed differences for AVP, VIP, or SS in *Six3<sup>syn</sup>* SCN as compared to control. However, it remains possible that altered AVP, VIP, or SS release patterns could contribute to changes in SCN network function and output in these mice. Impaired circadian time keeping is known to be a risk factor for obesity and metabolic dysfunction. SCN ablation causes increased insulin resistance, glucose intolerance, and weight gain [67]. But the relationship between circadian regulation and metabolism is probably complex. For instance, a high fat diet can blunt the normal circadian release of dopamine in the region around the SCN, and this may at least partly mediate its detrimental metabolic effects [67]. In our study, it is unclear how the weakened SCN output in *Six3<sup>syn</sup>* males might be related to the improved glucose tolerance and reduced fat mass, which is not what would be predicted based on the simple hypothesis that weakened SCN circadian output impairs metabolic function. Circadian effects on metabolism may depend on phase rather than simply strength of SCN circadian output. Indeed, V02 and VC02 phase mis-alignment in transgenic mice has previously been observed in mice lacking SCN VIP-VPAC2 signaling [51], where these mice presented a phase advance in their daily rhythms of metabolism and feeding [51]. Interestingly, this phase advance in feeding was associated with reduced metabolic rates. Although our data trended toward phase delays (V02, *p* = 0.162, *Figure 5O*), we did not find significant phase differences in the peak times of V02 or VC02 in *Six3<sup>syn</sup>* mice vs. controls.

Another possible explanation for the improved metabolic status despite apparently weakened SCN circadian output is the increased nocturnal activity of *Six3<sup>syn</sup>* males (observed in the metabolic cages, but not on the running wheels), which might actually indicate a selective strengthening of SCN rhythmic output. It is important to note two major differences between metabolic and wheel-running cages, which differed by size, and the absence/presence of a running wheel, respectively. The increase in nocturnal activity in the metabolic cages, was associated with increased repetitive behavior, which might indicate nocturnal neuronal hyperexcitability caused by increased repetitive behavior. Together, this increase in nocturnal activity could provide a strengthened metabolic signal to the body, overriding the weakened SCN circadian strength. Indeed, exercise is known to alter clock gene expression in muscle and several other peripheral tissues, and timed

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**Figure 7:** *Six3<sup>syn</sup>* males exhibit reduced GH release and reduced body length. A) Simplified schematic of the neuroendocrine axis regulating growth. B) Pituitary qPCR for Gh and C) circulating GH in adult *Six3<sup>fl/fl</sup>* and *Six3<sup>syn</sup>* 6- and 12-week-old males. D) Body length of *Six3<sup>fl/fl</sup>* and *Six3<sup>syn</sup>* males at 3- to 16 weeks of age. (B, C) Student’s t-test, and (D) Two-way ANOVA, repeated measures, *p* < 0.05, ***p* < 0.01, ****p* < 0.001, n = 4--9. Abbreviations: SS: somatostatin, GHRH: growth hormone releasing hormone, GH: growth hormone, IGF1: Insulin-like growth factor 1.
functioning reproductive axis, with normal hypothalamic expression of Six3syn release. This agrees with our previous report showing that Six3syn metabolism despite apparently weakened SCN circadian output in-potent circadian synchronizer [68].

Figure 8: Six3syn males exhibit reduced bone density and mineralization. A, B) qPCR for liver Ghr1 and Ghr in 12–16-week-old Six3fl/fl and Six3syn males. DEXA measurements of Six3fl/fl and Six3syn males fed standard chow revealed reduced C) bone mineral count, and D) bone mineral density. Student’s t-test, **p < 0.01, ***p < 0.001, n = 6. Abbreviations: Ghr: growth hormone receptor, Igf1: Insulin-like growth factor 1.

exercise combined with light therapy or food entrainment can be a potent circadian synchronizer [68–70]. Or perhaps the improved metabolism despite apparently weakened SCN circadian output involves the deletion of Six3 in the striatum of Six3syn mice. SIX3 is known to be involved in striatal medium spiny neuron plasticity and striatum function [27–29]. As the striatum is important for motor function and reward behavior, it would be of interest to specifically evaluate wheel-running patterns and metabolism in mice with Six3 deletion only in striatal medium spiny neurons.

4.3. Six3syn males have normal fertility

Despite their stunted growth and deregulation of the growth axis, Six3syn males had normally timed pubertal onset and a normally functioning reproductive axis, with normal hypothalamic expression of Gnrh1 supporting normal gonadotropin hormone production and release. This agrees with our previous report showing that Six3syn males generate a comparable number of litters to Six3fl/fl in a fertility assay [25]. The maintenance of fertility in Six3syn males is in striking contrast to the sub-fertility or infertility of female Six3syn mice, which is due to loss of SCN gating of the LH surge to the proper time of day, and consequent failure to ovulate [25]. Interestingly, we found a shorter free-running circadian period in the epididymis of Six3syn males, but this did not impact overall male fertility, so the possible function of local circadian clocks in this tissue is unclear. Circadian expression of core clock genes has also been found previously in other male reproductive tissues. In extra-testicular ducts and accessory tissues [71], the clock gene Bmal1 may regulate the production of testosterone [72], but this hormone is present at normal levels in Six3syn mice [25].

4.4. Value and limits of the Six3syn mouse model to study neuronal maturation and function

Using conditional KO mice (Cre-LoxP system) comes with numerous potential caveats to consider in data interpretation [43,73–78]. We have further validated the effectiveness of the Synapsincre allele for inducing recombination of the Six3syn allele throughout the brain, including the SCN, PVN, and PeVN [25]. Importantly, we did not observe any effects on brain peptide expression or behavioral measures in mice heterozygous for Synapsincre, providing reassurance that this allele on its own does not significantly alter neuronal function [25]. We here extend our previous study and show the Synapsincre allele recombines Six3syn in the male hypothalumus, cortex, pituitary, and testis. Of these tissues, Six3syn is known to be expressed in brain (hypothalamus and cortex), and pituitary. The identified recombination in the testis contrasts with our previous report, where we did not find Synapsincre allele recombination in the testis. This discrepancy could be due to the flox-allele recombined where the Rosa-allele used as a reporter for Synapsincre-driven recombination might be less efficiently recombined than the Six3syn allele, not allowing us to detect recombination in the testis in our previous study [25]. We do not know if Synapsincre targets somatotrophs, a cell population that expresses Six3 (personal correspondence with authors Ho et al., 2020) [79]. Although we cannot rule out the possibility that Six3 might have been deleted from a small number of somatotrophs, Six3syn males produced and secreted GH in a pattern consistent with the observed changes in SS. This indicates that somatotroph function was not substantially impaired in our Six3syn males.

Despite some limitations, the broad targeting of the Synapsincre allele in post-proliferative neurons provides a powerful tool to manipulate both maturing and mature neurons throughout the period of neuronal proliferation, which can dilute the effects of more conventional approaches. The specificity of targeting to later developmental stages is a particular advantage when studying genes, such as SIX3, that are highly expressed from early development through adulthood, and whose developmental loss can induce major brain defects obscuring any later effects. Thus, in this initial study, our aim was to obtain an overall picture of physiological functions of SIX3 in post-proliferative neurons, rather than to elucidate the precise roles of SIX3 within specific neuronal populations. For this broad objective, the broad targeting of Synapsincre in the brain provided an excellent tool to identify the novel roles of SIX3 in neurons.

4.5. Conclusion and future directions

Owing to broad expression of Synapsincre in the brain, allowing deletion of Six3 after neuronal proliferation, we have identified the novel roles of SIX3 in maturing and mature neurons, where we show it is required from ~4 weeks of age onward to regulate growth. Specifically, our data point to the novel role of SIX3 in the PeVN, where it appears to be a regulator of SS neuron numbers and metabolism. This study lays the foundation for future studies focusing more specifically on the mechanisms by which SIX3 acts within hypothalamic regions including PVN, PeVN, and SCN to regulate metabolic status and circadian rhythms.

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CONFLICT OF INTEREST

None declared.

APPENDIX A. SUPPLEMENTARY DATA

Supplementary data to this article can be found online at https://doi.org/10.1016/j.molmet.2021.101431.

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