Luciferase Reporter Mice for *In Vivo* Monitoring and *Ex Vivo* Assessment of Hypothalamic Signaling of **Socs3** Expression

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Suppressor of cytokine signaling-3 (SOCS3) is a negative regulator of actions of cytokines and the metabolic hormone leptin. In the hypothalamus, SOCS3 is induced in response to several conditions such as inflammation and high-fat diet feeding, modulates cellular signaling of cytokines and leptin, and mediates the effects of these biological conditions. However, signaling mechanisms controlling hypothalamic **Socs3** expression remains to be fully established. To facilitate the identification of molecular pathways of **Socs3** induction, we generated a real-time gene expression reporter mouse of **Socs3** (**Socs3**-Luc mice). We successfully detected a remarkable increase in luciferase activity in various tissues of **Socs3**-Luc mice in response to a peripheral injection of lipopolysaccharide, a potent inducer of inflammation, reflecting expression levels of endogenous **Socs3** mRNA. Using *ex vivo* hypothalamic explants of **Socs3**-Luc mice, we demonstrate that hypothalamic luciferase activity was significantly elevated in slices stimulated with known inducers of **Socs3** such as proinflammatory cytokines IL-6, IL-1β, and TNF-α, lipopolysaccharide, and cAMP-inducing agent forskolin. Using the *ex vivo* model, we found glycosyn synthase kinase-3 (GSK3)β-specific inhibitors to be potent inducers of **Socs3**. Furthermore, pharmacological inhibitors of β-catenin, a downstream mediator of GSK3β signaling, reduced **Socs3** luciferase activity *ex vivo*. Finally, hypothalamic inhibition of GSK3β hindered leptin-induced phosphorylation of signal transducers and activators of transcription 3 in hypothalamic explants. These results suggest that the **Socs3**-luciferase mouse is useful for *in vivo* monitoring of **Socs3** gene expression and for *ex vivo* slice-based screening to identify signaling pathways that control **Socs3** in the hypothalamus.

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**Freeform/Key Words:** **Socs3**, hypothalamus, luciferase reporter, cytokines, leptin

Suppressor of cytokine signaling-3 (SOCS3) is an inducible inhibitor of cytokine signaling [1, 2]. SOCS3 is induced by various cytokines including IL-6 family cytokines such as leptin and IL-6 [3]. Cytokines stimulate SOCS3 induction primarily via the Janus kinases (JAKs) and the signal transducers and activators of transcription (STATs) pathway. Induced SOCS3 inhibits cytokine signaling via the inhibition of JAK activity and targets the receptor complex for proteosomal degradation [4]. Thus, SOCS3 is thought to limit the signaling cascade to prevent an excessive, aberrant response of cytokines.

Abbreviations: BAC, bacterial artificial chromosome; DAB, 3,3′-diaminobenzidine; GSK3, glycosyn synthase kinase-3; JAK, Janus kinases; LepR, leptin receptor; LPS, lipopolysaccharide; Luc, Luciferase; MSAB, methyl 3-4methylphenyl sulfonyl amino benzoate; NF-κB, nuclear factor κB; **Socs3**, suppressor of cytokine signaling-3; STAT3, signal transducers and activators of transcription 3; WT, wild type.
SOCS3 in the hypothalamus has emerged as a critical regulator of energy balance [4–8]. SOCS3 is significantly induced in the hypothalamus in response to high-fat diet feeding, pregnancy, aging, and long-day photoperiod that promotes positive energy balance [5, 9]. Forced expression of SOCS3 in a distinct population of hypothalamic neurons promoted body weight gain and adiposity [10]. Consistently, induction of the constitutively active form of STAT3 in pro-opiomelanocortin neurons stimulated Socs3 expression and increased body weight [11]. In contrast, brain-specific Socs3 knockout or its haploinsufficiency mice were significantly protected against the development of diet-induced obesity [6, 7]. Furthermore, the range of hypothalamic neuron-specific Socs3 deletion mouse models demonstrates that hypothalamic Socs3 critically regulates body weight and glucose homeostasis [12–14]. Thus, the level of Socs3 expression in the hypothalamus seems to determine body weight and adiposity in vivo.

The effects of hypothalamic Socs3 on energy homeostasis seem to be at least in part mediated through inhibiting leptin actions. Leptin rapidly induces Socs3 expression in the hypothalamus [15]. Increased Socs3 expression leads to the attenuation of cellular signaling of leptin in cultured cells and in animals in a cell-autonomous manner [10, 16, 17]. In contrast, Socs3 deficiency enhances cellular and physiological responses to exogenous leptin [6, 7]. Thus, it is well established that Socs3 is a negative feedback inhibitor of leptin in vivo. Signaling mechanisms mediating leptin-induced Socs3 involve the JAK-STAT3 pathway [15, 18–20]. In addition, recent studies have suggested the existence of a leptin-STAT3-independent mechanism. For example, STAT3 deficiency within leptin receptor (LepR)-expressing hypothalamic cells did not decrease mRNA expression of Socs3 in LepR cells [21]. Along the same line, inactivation of Socs3 in LepR cells displayed a minimum effect on body weight [14]. An increase of Socs3 in LepR neurons does not cause obesity [10]. Clearly, additional work is needed to elucidate signaling networks responsible for Socs3 expression.

Socs3 is also induced by other signaling pathways. Socs3 expression is increased in response to stimulation with inflammatory signals such as Toll-like receptor 4 [22], TNF-α [23], and IL-1β [23] that commonly activate the inhibitor of κ B kinase/nuclear factor κB (NF-κB) pathway [2, 24]. Indeed, activation of the NF-κB pathway is sufficient to induce Socs3 in the hypothalamus and elicit hypothalamic resistance to exogenous leptin [17]. In addition, numerous studies demonstrate that cAMP plays a role to downregulate IL-6 signaling as well as to induce Socs3 in various cell types [25–30]. Importantly, cAMP-elevating agents were reported to diminish cellular sensitivity to exogenous leptin in endothelial cells [30] and in the hypothalamus [31]. These findings raise a possibility that a molecular pathway of Socs3 induction might also interfere with cellular leptin actions. However, the signaling networks that confer the control of Socs3 levels and leptin sensitivity remain to be fully established. Therefore, delineation of molecular pathways responsible for Socs3 induction is of high interest. To facilitate the identification of such molecular pathways, we generated a new Socs3-luciferase reporter mouse (Socs3-Luc) and used hypothalamic explants of Socs3-Luc to search for and characterize a signaling pathway that influences Socs3 expression and cellular leptin action ex vivo.

1. Materials and Methods

A. Generation of Socs3-Luc Mice

A 98.6-kb bacterial artificial chromosome (BAC; bMQ-126K3; Source BioScience) contains the entire Socs3 gene locus. Luciferase2 cDNA was prepared from pGL4.10 (Promega, Madison, WI). The luciferase2 cDNA was introduced into the translation start site of the Socs3 gene in the BAC clone in Escherichia coli cells using BAC recombineering techniques [32]. The recombinant Socs3 luciferase (Socs3-Luc) BAC DNA was prepared by NucleoBond Xtra BAC Kit (Macherey-Nagel, Germany), linearized by PI-SceI digestion, purified with phenol-chloroform, and precipitated with ethanol. Transgenic mice were obtained by injecting the linearized Socs3-luciferase BAC DNA into pronuclei of C57BL/6 mice in the Genetically Engineered Mouse Core at Baylor College of Medicine. Socs3-Luc founders were maintained at a C57BL/6 background. Tail biopsies were used for the genotyping on PCR with primers TL luciferase-F6 5’-CTTCGACGACATGATAAGATC-3’ and
MFSOCS3-F1 5’-TGACGCTCAACGTGAAGAAG-3’ producing the amplicon of 591-bp long. The Institutional Animal Care and Use Committee at Baylor College of Medicine approved all procedures to maintain and use these mice.

B. Protein Extraction and Luciferase Assay

*In vitro* luciferase activity was quantified using the Luciferase Assay System (Promega, Madison, WI). Tissues were extracted using the 1× Luciferase Cell Culture Lysis Reagent (Promega, Madison, WI), sonicated and vortexed, followed by centrifugation for 10 minutes at 4°C and 15,000 rpm. Supernatant was placed in a new 1.5-mL tube and stored at −80°C. To assess luciferase activity, 4 μL of sample was mixed with 20 μL Luciferase assay reagent, and luminescence was measured using a luminometer (Promega, Madison, WI). The Promega Luc-0-INJ protocol was used to measure luciferase activity in the tissues.

C. In Vivo Imaging

Whole body imaging of luciferase activity *in vivo* was monitored using an In Vivo MS FX PRO imaging system (Bruker, Billerica, MA). Male and female mice were injected IP with 150 mg/kg body weight of D-luciferin solution (Promega, Madison, WI) and anesthetized (2.5% isoflurane). Mice were transferred to nose cones within the chamber and imaged 10 minutes after injection of D-luciferin. Exposure time was 10 seconds on the ventral side, and luminescence was quantified using the Bruker imaging software. Bright-field images were also taken of mice with an exposure time of 5 seconds. Tissues, collected from male mice, were also imaged using the In Vivo MS FX PRO. Briefly, male mice were injected IP with 150 mg/kg body weight of D-luciferin solution and dissected 10 minutes later. Tissues were placed in chamber and bright-field, and luminescence images were taken using the same exposure times for *in vivo* imaging.

D. Organotypic Slice Experiments

Screening of compounds was accomplished using an organotypic slice culture system. Hypothalamic slices were made essentially as described before [31]. Briefly, C57BL/6 mice male and female pups, 8 to 12 days old, were decapitated, and the brains were quickly removed. Hypothalamic tissues were blocked and sectioned in depth of 250 μm on a vibratome (VT1000S, Leica) in chilled Gey’s Balanced Salt Solution (Invitrogen, Carlsbad, CA) enriched with glucose (0.5%) and KCl (30 mM). The coronal slices containing the arcuate nucleus were then placed on Millicell-CM filters (Millipore, pore size 0.4 μm, diameter 30 mm), and maintained at an air-media interface in MEM (Invitrogen, Carlsbad, CA) supplemented with heat inactivated horse serum (25%, Invitrogen, Carlsbad, CA), glucose (32 mM) GlutaMAX (2 mM, Invitrogen, Carlsbad, CA). Cultures were typically maintained for 10 days in standard medium, which was replaced three times a week. After 10 days, the slices were used for experiments. The following modifications were made for *Socs3*-luc slices: after brains were sectioned, all sections were cut to the same diameter before being placed on Millicell-CM filters in culture. Slices were maintained 1 to 7 days in standard medium before being transferred to individual cell culture petri dishes (Thermo Fisher Scientific, Waltham, MA) containing low-serum (2% heat inactivated horse serum) phenol-red free MEM supplemented with glucose (32 mM) GlutaMAX (2 mM), 10% HEPES buffer (Invitrogen, Carlsbad, CA), and 0.1 mM D-luciferin. Dishes were sealed with a circular cover glass (Thermo Fisher Scientific, Waltham, MA) using vacuum grease (VWR, Radnor, PA) and placed in a Lumicycle 32 (Actimetrics, Wilmette, IL), a 32-channel luminometer at 37°C to measure luminescence from tissues. Treatments were added 2 days after slices were placed in the Lumicycle. Luciferase activity was analyzed using Lumicycle Analysis software.

E. Reagents

We used the following reagents: lipopolysaccharide (LPS; L4391, Sigma-Aldrich, St. Louis, MO); IL-6 (RMIL6I, Thermo Fisher Scientific, Waltham, MA), IL-1β (I5271, Sigma-Aldrich St. Louis,
MO); TNF-α (RMTNFAI, Thermo Fisher Scientific, Waltham, MA); forskolin (#1099, TOCRIS, Bristol, United Kingdom); LY2090314 (SML1438, Sigma-Aldrich); CHIR99021 (SML1046, Sigma-Aldrich St. Louis, MO); methyl 3-4methylphenyl sulfonyl amino benzoate (MSAB; M60316, Xcess Biosciences, San Diego, CA); and FH535 (S7484, SelleckChem, Houston, TX).

**F. Quantitative RT-PCR**

Tissues were quickly removed and frozen in liquid nitrogen and kept in −80°C until further processing. Total RNA was extracted using the RNeasy Lipid Tissue Kit (QIAGEN Sciences, Germantown, MD). cDNA was generated by iScript RT Supermix (Bio-Rad Laboratories, Hercules, CA) and used with SsoAdvanced Universal SYBR Green Supermix (Bio-Rad Laboratories, Hercules, CA) for quantitative real-time PCR analysis. Quantitative PCR assays were performed using a CFX384 Touch Real-Time PCR Detection System (Bio-Rad Laboratories, Hercules, CA). Normalized mRNA levels were expressed in arbitrary units obtained by dividing the averaged, efficiency-corrected values for sample mRNA expression by that of 18SrRNA expression for each sample. The resulting values were expressed as fold change above average control levels. The primer sequences are as follows: SOCS3 (F-CACCTGGACTCCTATGA-GAAAGTG and R-GAGCATCATACTGATCCAGGAACT), PTP1B (F-GGAACAGGTACCGAGATGTCA and R-AGTCATTATCTTCCTGATGCAATT), TCPTP (F-AGGGCTTCCTTCTAAGG and R-GTTTCATCTCTGCTGACCTTTCTGAG), and 18SrRNA (F- CACGGACAGGATTGACAGATT and R-GCCAGAGTCTCGTTCGTTATC).

**G. Immunohistochemistry**

For organotypic slices, slices were fixed overnight in 4% paraformaldehyde at 4°C. Then slices were cut out from the membrane and rinsed three times for 10 minutes each in PBS, pH 7.4 and then for 20 minutes in 1% hydrogen peroxide and 1% sodium hydroxide in PBS to quench endogenous peroxidase activity. Following a series of washes with PBS, slices were incubated for 48 to 72 hours at 4°C in pSTAT3 antibodies (1: 3000, 9131, Cell Signaling Technology, Boston, MA) [33] in 3% normal donkey serum (Jackson ImmunoResearch Laboratories, West Grove, PA) with 0.25% Triton X-100 in PBS and 0.02% sodium azide. After washing in PBS, slices were incubated in a biotinylated goat antirabbit antibody IgG (1:1000, BA-1000, Jackson ImmunoResearch Laboratories West Grove, PA) [34] in 3% donkey serum in 0.25 % Triton X-100 in PBS for 1 hour at room temperature. Tissues were then rinsed in PBS and incubated in the avidin-biotin-peroxidase complex kit (1:500, Vectastain Elite ABC kit; Vector Laboratories, Burlingame, CA) for 1 hour. Slices were washed in PBS then reacted in 3,3’-diaminobenzidine (DAB; Sigma-Aldrich St. Louis, MO). Slices were then rinsed in PBS and mounted on slides using Vectashield (Vector Laboratories Burlingame, CA). For fluorescence staining, slices were incubated with Streptavidin, Alexa Fluor™ 488 conjugate (Molecular Probes, Eugene, OR). Immunohistochemical images were analyzed using a bright-field Leica microscope. To measure pSTAT3 signal in the arcuate nucleus, uneven background was eliminated with Adobe Photoshop and then intensity of DAB staining was measured.

**H. Statistics**

The data are presented as mean ± SEM. Statistical analyses were performed using GraphPad Prism for a two-tailed unpaired Student t test or one- or two-way ANOVA followed by post hoc Tukey, Bonferroni, or Sidak tests. P < 0.05 was considered to be statistically significant.

**2. Results**

**A. Generation and Characterization of Socs3 Luciferase Reporter Mice**

To create an in vivo monitoring system for endogenous Socs3 expression in a noninvasive manner, we generated transgenic mice (Socs3-Luc) by engineering a Socs3 BAC such that
Luciferase2 gene is driven by Socs3 regulatory elements (Fig. 1A–1C). The firefly luciferase-polyA was inserted into the ATG site of the Socs3 gene in a 93.6-kb BAC containing the entire Socs3 gene locus via homologous recombination, and transgenic mice were generated by pronuclear injection of the modified BAC. Transgenic founders and their offspring were identified by PCR (Fig. 1B). With the use of an in vivo imaging system, we functionally validated Socs3-Luc transgenic mice by monitoring luciferase activity in the reporter mice. In vivo luciferase imaging detected robust luciferase signals in Socs3-Luc but not in wild type (WT) mice (Fig. 1C). Analyzing luciferase activity of tissue lysates from liver and hypothalamus known to express Socs3 we further confirmed that luciferase activity, normalized to the amount of total protein, was specifically detected only in tissues of Socs3-Luc mice (Fig. 1D).

**Figure 1.** Generation of Socs3-Luc mice. (A) Schematic of the Socs3-Luc transgene. The full-length luciferase gene was inserted into the ATG codon of the Socs3 gene in a BAC (bMQ126k03). (B) Socs3-Luc mice were identified by PCR analysis. (C) Live imaging of luciferase activity in WT and Socs3-Luc mice. (D) Luciferase enzymatic activity in liver and hypothalamus of WT and Socs3-Luc mice.
Next we examined whether the reporter mice respond to LPS endotoxin [22] that activates a potent, common proinflammatory response and is known to induce Socs3 [2]. We injected LPS into the periphery of Socs3-Luc mice and their littermate controls (WT) and measured luciferase activity using the in vivo imaging system. Total body luciferase activity was markedly stimulated in the Socs3-Luc mice 4 hours after LPS administration, whereas luciferase activity was low in vehicle-injected animals (Fig. 2A). To assess the tissue distribution of luciferase luminescence detected in Socs3-Luc, we dissected internal peripheral organs including small intestine, liver, spleen, kidney, stomach, pancreas, white adipose tissue, brown adipose tissue and heart, at 4 hours after the LPS administration and measured their luciferase activity using in vivo imaging system analysis. Almost all of the tissue samples showed some level of luciferase luminescence induced by LPS (Fig. 2B). We also determined if the Socs3 luciferase reporter recapitulates LPS-induced gene expression pattern of Socs3 mRNA by measuring in vitro biochemical luciferase activity and endogenous Socs3 mRNA in various tissues of Socs3-Luc mice with or without LPS injection. The basal levels of luciferase activities in tissue lysates were detected in all tissues we examined with relatively high in vitro luciferase activity from liver, stomach, muscle, and spleen (Fig. 2C). Upon LPS treatment, the highest induction of luciferase activity was found in intestine, pancreas, kidney, and liver, whereas it was moderate in brain tissues, fat tissues, stomach, and muscle (Fig. 2D). We found a close correlation between luciferase activity and endogenous Socs3 mRNA expression (Fig. 2D and 2E). Endogenous Socs3 mRNA expression was remarkably induced in most of the tissues in an LPS-dependent manner (Fig. 2E). In particular, liver, kidney, stomach, heart, intestine, and pancreas showed highly induced expression of the endogenous Socs3 (Fig. 2E), which closely correlated with the induced expression pattern of the luciferase luminescence. These data thus demonstrate that the Socs3 reporter model faithfully mirrors the tissue distribution of endogenous Socs3 mRNA in vivo.

B. An Ex Vivo Model to Monitor Hypothalamic Socs3 Expression

Hypothalamic levels of Socs3 crucially regulate energy balance [5, 35]. To explore signaling mechanisms by which hypothalamic Socs3 levels are determined, we used hypothalamic explants of Socs3-Luc mice that allow us to pharmacologically manipulate various signaling cascades and rapidly assess its effect on Socs3 expression. Mediobasal hypothalamus was dissected and placed on a membrane in a dish containing media supplemented with luciferin substrate. Hypothalamic explants were treated with chemicals and peptides, and luminescence was continuously measured in real time with photomultiplier tube detectors during the treatment (Fig. 3A). To confirm that the Socs3 luciferase reporter hypothalamic slice is capable of responding to LPS, we treated slices with LPS and luminescence was continuously measured in real time. As shown in Fig. 3B, slices treated with LPS show a 3.0-fold increase in luciferase activity compared with their PBS controls. Because Socs3 is also produced by proinflammatory cytokines that include IL-6, TNF-α, and IL-1β [22, 23], slices were treated with them to further validate our ex vivo model. As expected, all three of the cytokines tested resulted in an increase in luciferase activity (Fig. 3C–3E). Forskolin, a previously reported inducer of Socs3 in various cells, also displayed a 2.4-fold increase in luciferase activity in hypothalamic explants (Fig. 3F). These results confirm that hypothalamic slices prepared from Socs3-Luc mice can serve as a suitable screening tool for measuring the effects of modulators of Socs3 expression in the hypothalamus.

C. Inhibitors of Glycogen Synthase Kinase-3β Induces Socs3 Gene Expression in the Hypothalamus

With the use of the ex vivo model to screen compounds for their ability to alter Socs3 expression, we came upon chemicals that target proteins involved in the glycogen synthase kinase-3 (GSK3)β-β-catenin signaling pathway. First, we found that treatment with GSK3 inhibitor LY2090314 [36], a potent and selective small-molecule inhibitor of GSK3α and
Figure 2. Functional validation of Socs3-Luc mice. (A) Luminescence images of Socs3-Luc mice intraperitoneally administered with either saline or 100 ng/kg LPS for 4 h. (B) Bright field images (left) and bioluminescence images (right) of various tissues of Socs3-Luc mice receiving LPS (100 ng/kg for 4 h) or saline. (C and D) In vitro luciferase activity of the indicated tissue lysates extracted from mice treated with either (C) saline or (D) 100 ng/kg LPS for 4 h shown as bioluminescent signal. The same amount of protein (3.5 µg) was used for each reaction. (E) Socs3 mRNA in various tissues of the mice treated with either saline or 100 ng/kg LPS for 4 h. Data are mean ± SEM; n = 3 to 4/per group. *P < 0.05; **P < 0.01; ****P < 0.0001 vs control based on t tests.
GSK3β isoforms, induces a 2.1-fold increase in luciferase activity and this occurs in a dose-dependent manner (Fig. 4A and 4B). Consistently, the mRNA level of endogenous Socs3 was elevated in slices treated with LY2090314 (Fig. 4C). Interestingly, we did not detect any increases in other inhibitors of cytokine signaling such as tyrosine protein phosphatases. Hypothalamic induction of Socs3 was also observed in explants treated with CHIR99021 [37], a well-established specific inhibitor for GSK3β with a different kinase selectivity profile than LY2090314 (Fig. 4D and 4F). This increase also occurred in a dose-dependent manner (Fig. 4E). These data clearly suggest that pharmacologic inhibition of GSK3β stimulates Socs3 gene expression in the hypothalamus.

D. Inhibitors of β-Catenin Decrease Socs3 Gene Expression in the Hypothalamus

Next, we explored the mechanism by which GSK3β inhibition increased Socs3 transcription. Because GSK3β acts through β-catenin [38], we hypothesized that the action of GSK3β inhibitors in hypothalamus could be mediated through β-catenin. To test this, Socs3-Luc slices were treated with FH535 (50 μM) [39], an inhibitor of β-catenin. This treatment resulted in a decrease in luciferase activity in a dose-dependent manner (Fig. 5A and 5B). Because FH535 is not a selective inhibitor of β-catenin [39], we further tested a more specific β-catenin inhibitor MSAB [40] for its ability to decrease Socs3 luciferase signal. Similar to FH535, treating slices with MSAB significantly decreased Socs3 luciferase activity, and this
decrease was also dose-dependent (Fig. 5C and 5D). These data suggest that inhibition of β-Catenin induces Socs3 gene expression.

E. A GSK3β Inhibitor Blocks Leptin-Induced STAT3 Phosphorylation in Hypothalamic Slices

SOCS3 potently limits cellular leptin signaling in the hypothalamus. To examine the functional consequences of GSK3β inhibition, we examined if the treatment desensitizes the hypothalamic cells to leptin using ex vivo hypothalamic slices that recapitulate hypothalamic leptin signaling [31]. Hypothalamic slices were exposed to CHIR99021 (10 μM for 6 hours) followed by leptin stimulation (120 nM for 60 minutes) to examine the effects of GSK3β inhibition. Similar to previous observations, leptin robustly induced phosphorylation of STAT3 in control slices. In contrast, treatment of the slices with both CHIR99021 had a potent inhibitory effect on leptin-induced STAT3 phosphorylation in the hypothalamus (Fig. 6A and 6B). CHIR99021 alone had no effect on pSTAT3 (Fig. 6A and 6B). These data suggest that inhibition of GSK3β impairs hypothalamic LepR signaling.

3. Discussion

SOCS3 plays a critical role in controlling inflammation and metabolism. SOCS3 levels appear to determine cellular sensitivity to cytokines and metabolic hormones, and SOCS3
gene expression is tightly controlled. However, the underlying signaling mechanisms are not fully established. To explore the mechanism of \textit{Socs3} induction, we have generated \textit{Socs3} reporter mice that allow us to quantitatively monitor changes in \textit{Socs3} expression \textit{in vivo}. We have also developed a hypothalamic slice-based compound screening platform to monitor hypothalamic \textit{Socs3} expression. The tool provides an opportunity to identify signaling pathways that control \textit{Socs3} expression in hypothalamic \textit{ex vivo}.

Using the LPS-induced inflammation model that stimulates \textit{Socs3} induction \textit{in vivo} [1, 23, 41], we validated if \textit{Socs3}-Luc mice could recapitulate the LPS-induced robust induction of \textit{Socs3}. \textit{Socs3} luciferase activity was detected in almost all tissues among those examined after the LPS injection. Importantly, the tissue distribution pattern of LPS-induced luciferase
signals closely correlates with that of endogenous Socs3 mRNA. In accordance with the reported studies showing that Socs3 induction emerges only after a few hours following the LPS injection in vivo [42, 43], we found that luciferase signals were observed at 4 hours, but not at 30 minutes after LPS injection in the Socs3-Luc transgenic mice (data not shown). This pattern of luciferase signals was observed in both whole-body mice and individual tissues we examined (small intestine, liver, spleen, kidney, stomach, pancreas, WAT, BAT, and heart). These results are consistent with previous studies that have shown peak expression levels of Socs3 to be around 4 hours in liver and brain tissues [42, 43]. A number of studies have shown that LPS increase SOSC3 rapidly within several minutes in cells such as macrophages in vitro and in vivo. Although the exact reasons for this discrepancy remain unknown, this might be due to difference in the time frame of temporal response to LPS in different tissues. These observations underscore that the Socs3 luciferase reporter model faithfully and robustly reports spatiotemporal dynamics of endogenous Socs3 mRNA in vivo.

To establish a reliable in vitro monitoring system, we first developed a hypothalamic slice culture-based assay that allows real-time monitoring of Socs3 expression using organotypic hypothalamic slices. The organotypic brain slice model is known to partly preserve the in vivo anatomy and function of the hypothalamus [44]. Previous results from the studies by ours and other groups demonstrated that the slice model can genuinely recapitulate in vivo cellular responses to metabolic hormones, anatomically, biochemically, and electrophysiologically [31, 45–48]. In the current study, we show that the ex vivo system reliably reproduces hypothalamic responses to LPS, TNF-α, IL-1β, IL-6, and forskolin, which increase Socs3 mRNA in vivo [22, 23, 26–30]. All of the Socs3 inducers remarkably elevated luciferase activity in Socs3-Luc hypothalamic explants. Thus, the in vivo and ex vivo data collectively suggest that luciferase activity serves as a genuine reporter for monitoring Socs3 expression in real-time, and more importantly, this slice-based imaging model offers a feasible and reliable platform to search for small chemicals for Socs3 expression in vitro.

With the use of the ex vivo monitoring model, we identified several compounds (CHIR99021, LY2090314, FH535, and MSAB) that lead to the modulation of Socs3 expression in the hypothalamus. LY2090314 [36], a potent and selective small-molecule inhibitor of GSK3α and GSK3β isoforms, clearly stimulated Socs3 luciferase activity in the
hypothalamic explants, suggesting that inhibition of GSK3 activity results in increased Socs3 gene expression. Further, hypothalamic induction of Socs3 was observed in explants treated with CHIR99021 [37], a structurally distinct specific inhibitor for GSK3\(\beta\). The effects of LY2090314 and CHIR99021 were observed in the nanomolar or micromolar range, respectively, which are within the effective concentrations established by various cell-based functional assays [36, 49, 50]. GSK3\(\beta\) is known to be involved in several distinct signaling pathways such as cellular signaling mediating insulin, wingless and integration site growth factor, and hedgehog. GSK3\(\beta\) is constitutively active under unstimulated conditions where GSK3\(\beta\) phosphorylates \(\beta\)-catenin, which in turn is subjected to ubiquitin proteasome degradation. In contrast, inhibition of GSK3\(\beta\) by upstream signals such as wingless and integration site growth factor and insulin signaling results in an increase in \(\beta\)-catenin protein levels, which promotes the transcription of its target genes in a T-cell factor/lymphoid enhancer factor-dependent mechanism. Thus, pharmacological inhibition of GSK3\(\beta\) should also lead to activation of \(\beta\)-catenin-dependent gene expression. \textit{Ex vivo} studies suggest that GSK3\(\beta\) inhibition-induced Socs3 expression is mediated by \(\beta\)-catenin (data not shown), indicating a pathway that links GSK3\(\beta\) and Socs3 via \(\beta\)-catenin in the hypothalamus. The findings in the \textit{ex vivo} studies presented in this paper provide evidence demonstrating that the GSK3\(\beta\)–\(\beta\)-catenin pathway might act as a potential regulator of Socs3 expression in the hypothalamus.

One potential implication of our results is that GSK3\(\beta\) inhibition may downregulate cellular signaling of leptin via Socs3 induction. Several studies have shown that Socs3 inhibits cellular actions of leptin by interfering with LepR signaling. Prior studies demonstrated that Socs3 is induced by several distinct signaling pathways that include the JAK-STAT3 pathway, the inhibitor of \(\kappa\) B kinase/NF-\(\kappa\)B pathway [2, 17, 24], the endoplasmic reticulum stress-related pathways [51], and the cAMP-related pathways [30, 31]. Notably, all of these pathways limit cellular leptin sensitivity, as well as induce Socs3 in the hypothalamus implicating that the pathways underlying Socs3 induction would inhibit cellular signaling of leptin. Consistent with this notion, our studies indeed demonstrate that a specific inhibitor for GSK3\(\beta\) inhibited leptin-induced STAT3 phosphorylation, while inducing hypothalamic Socs3. The failure of exogenous leptin to modulate key signaling events such as STAT3 phosphorylation within hypothalamic neurons is well documented in various physiological and pathophysiological settings. These phenomena occur in response to high-fat diet feeding, chronic leptin exposure, pregnancy and lactation, age, and seasonal variations in the length of the photoperiod in seasonal animals. Under these conditions, Socs3 is commonly elevated and likely to play a role in suppressing leptin actions. Future studies are warranted to elucidate the role of the GSK3\(\beta\)–\(\beta\)-catenin pathway in Socs3 induction under these conditions.

In addition to leptin actions, SOCS3 commonly suppresses cellular signaling of inflammation. A number of studies have shown that inhibition of GSK3\(\beta\) has the ability to reduce inflammation. For example, GSK3\(\beta\) inhibition protects against LPS-induced endotoxin shock \textit{in vivo}, decreases proinflammatory cytokine production, and interferes with NF-\(\kappa\)B and STAT3 pathways. At the molecular level, GSK3\(\beta\) inhibition was reported to affect the nuclear amounts of transcription factors NF-\(\kappa\)B subunit p65 and CREB interacting with the coactivator CBP [52]. GSK3\(\beta\) was also shown to block STAT3 DNA binding activity and STAT3-dependent gene expression [53]. These data suggest the ability of GSK3\(\beta\) inactivation to strongly induce Socs3 and may add another mechanism by which GSK3 inactivation potently suppresses inflammation.

Because Socs3 expression is altered dynamically under various biological conditions such as inflammation, dietary obesity, pregnancy and lactation, seasonal adiposity, intestinal bowel disease, ulcerative colitis, Crohn disease, rheumatoid arthritis, thymocyte differentiation, and atherosclerotic lesions [5, 35, 54–58], this model may also facilitate the functional investigation of Socs3 in multiple tissues and in diverse physiological and pathophysiological conditions.

In summary, we have created a BAC-transgenic luciferase reporter mouse carrying the Socs3-luciferase reporter in the Socs3 genomic locus. These mice show robust Socs3 activity
after administration of LPS in vivo, and hypothalamic explants derived from these animals can be used as a slice-based screening platform to identify signaling networks that alter hypothalamic Socs3 expression.

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