Steroidogenic Factor 1 Regulates Expression of the Cannabinoid Receptor 1 in the Ventromedial Hypothalamic Nucleus

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The nuclear receptor steroidogenic factor 1 (SF-1) plays essential roles in the development and function of the ventromedial hypothalamic nucleus (VMH). Considerable evidence links the VMH and SF-1 with the regulation of energy homeostasis. Here, we demonstrate that SF-1 colocalizes in VMH neurons with the cannabinoid receptor 1 (CB1R) and that a specific CB1R agonist modulates electrical activity of SF-1 neurons in hypothalamic slice preparations. We further show that SF-1 directly regulates CB1R gene expression via a SF-1-responsive element at −101 in its 5′-flanking region. Finally, we show that knockout mice with selective inactivation of SF-1 in the brain have decreased expression of CB1R in the region of the VMH and exhibit a blunted response to systemically administered CB1R agonists. These studies suggest that SF-1 directly regulates the expression of CB1R, which has been implicated in the regulation of energy homeostasis and anxiety-like behavior.

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Abbreviations: ACEA, Arachidonyl-2′-chloroethylamide hydrate; aCSF, artificial cerebrospinal fluid; ANA, anandamide; BDNF, brain-derived neurotrophic factor; CB1R, cannabinoid receptor 1; CNS, central nervous system; DMH, dorsomedial hypothalamic nucleus; eGFP, enhanced green fluorescent protein; FBS, fetal bovine serum; ISH, in situ hybridization; JNK, c-Jun N-terminal kinase; KO, knockout; MA, methanandamide; NPY, neuropeptide Y; p, phosphorylated; SF-1, steroidogenic factor 1; STAT, signal transducer and activator of transcription; VMH, ventromedial hypothalamic nucleus; WT, wild type.

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RESULTS

SF-1 Is Required for CB1R Expression in the VMH

We used CNS-specific SF-1 KO mice to explore potential regulatory roles of SF-1 in CB1R expression in the VMH. As shown in Fig. 1, the effects of this conditional inactivation to abolish SF-1 immunoreactivity and perturb VMH structure (Fig. 1A) were consistent with our previous findings (7). As described (16, 17), CB1R transcripts in wild-type (WT) mice were expressed in many brain regions, including the cerebral cortex, amygdala, hippocampus, and hypothalamus (Fig. 1, B and C). CB1R expression was generally preserved in the CNS-specific SF-1 KO mice (Fig. 1B) but was decreased considerably in the region where the VMH normally resides (Fig. 1, B and C). These results indicate that the CNS-specific KO of SF-1 selectively impairs CB1R expression in the VMH, suggesting that it may be a direct target of SF-1 in the VMH neurons.

Electrophysiological Regulation of SF-1 Neurons by CB1R

To demonstrate colocalization of CB1R and SF-1 in VMH neurons, we first used quantitative RT-PCR assays with RNA samples prepared from neurons that express a SF-1-enhanced green fluorescent protein (eGFP) transgene (18). CB1R was expressed in these SF-1 neurons, and the level of CB1R expression was decreased by more than 50% in eGFP-positive cells isolated from SF-1 KO mice (Fig. 1D). These studies demonstrate that SF-1 neurons express CB1R and further suggest that SF-1 may regulate their expression of CB1R. The finding that some CB1R expression persists in the absence of SF-1 suggests that other transcription factors also activate CB1R expression in VMH neurons; it further suggests that the decreased levels of CB1R transcripts detected by in situ hybridization (ISH) (Fig. 1) may reflect both decreased transcript levels and greater dispersion of SF-1 neurons within the mediobasal hypothalamus, as previously described (6).

To explore possible functional effects of cannabinoids signaling in SF-1 neurons, we used the same SF-1/eGFP transgene described above to mark SF-1-positive and SF-1-negative VMH neurons. As described in Materials and Methods, presynaptic influences were eliminated by preincubation with a cocktail of inhibitors of glutamate, GABA_A, and glycine receptors and whole-cell patch-clamp recordings were collected in slice preparations that included the VMH. The average action potential frequency in SF-1 neurons was 6.2 ± 1 Hz (Fig. 2).

We first examined the direct effect of cannabinoids on the firing rate of these SF-1/eGFP-positive neurons. After at least 10 min of stable recording of action potentials, the CB1R receptor agonist WIN 55212-2 (1 μM) was added to the bath medium. WIN 55212-2 reduced the frequency of action potentials in approximately 60% of SF-1/eGFP-positive neurons tested (9 of 14 neurons; Fig. 2A). In the affected neurons, treatment with WIN 55212-2 significantly decreased the firing rate of SF-1 neurons from 6.2 ± 1 Hz to 2.6 ± 0.8 Hz (Fig. 2C), which was associated with a statistically insignificant hyperpolarization of membrane potential (control: −45 ± 1.5 mV, plus WIN: −47 ± 1.7 mV; n = 9). The effect of WIN 55212-2 to decrease excitability of SF-1 neurons was blocked by the CB1R antagonist, AM251, suggesting that the effect is specific for CB1R (data not shown). In contrast to the direct effect of cannabinoids on SF-1 neurons, the action potential frequency of SF-1/eGFP-negative VMH neurons in our slice preparations was not significantly altered by treatment with WIN 55212-2 (firing rate: control: 5.2 ± 2 Hz, plus WIN 55212-2: 4.8 ± 2 Hz; membrane potential: control, −48 ± 2.5 mV; plus WIN, −43 ± 1.8 mV; Fig. 2, B and C). These results thus show that functional CB1R is expressed in SF-1 neurons whose activity can be inhibited by a CB1R agonist.

Based on the effects of the SF-1 KO on CB1R expression in the VMH and of the CB1R agonist on action potential frequency of SF-1 neurons, we examined the impact of the CNS-specific SF-1 KO—and the resulting perturbation of VMH structure and CB1R expression—on the in vivo response to systemic administration of drugs that affect the cannabinoid system. Because the effects of cannabinoids on food intake involve hypothalamic action (3) and the VMH is a prominent site of CB1R expression (16, 19), we examined CB1 agonist and/or antagonist effects on nocturnal food intake in SF-1nCre;F/+ (WT) and SF-1nCre;F/- (KO) mice. After 24 h of fasting, mice were systemically administered either of two different CB1R agonists, methanandamide (MA) and arachidonyl-2′-chlo-roethylamide hydrate (ACEA), or the antagonist AM251, and then nocturnal food intake was measured over 3 h as described in Materials and Methods. Consistent with a previous report that rats given MA ate significantly more food in a partial satiety state (20), administration of MA or ACEA to WT mice significantly increased food intake (Fig. 3, A and B). This increased food intake was significantly diminished in CNS-specific SF-1 KO mice (Fig. 3, A and B). Consistent with the apparent preservation of VMH structure and SF-1 immunoreactivity (Fig. 1), mice heterozygous for the SF-1 KO (SF-1nCre+/−) responded to treatment with the CB1R agonists in a manner indistinguishable from WT mice (Kim, K., unpublished observation).

The CB1R antagonist AM251 significantly inhibited food intake in WT mice, but its anorexic effect was
significantly reduced in CNS-specific SF-1 KO mice relative to WT mice (Fig. 3C). Taken together, these results argue that SF-1 regulation of CB1R expression in the VMH plays an important role in cannabinoid-induced effects on food intake.

SF-1 Directly Regulates CB1R Promoter Activity

We next asked whether CB1R is a direct target of SF-1 transcriptional regulation. Inspection of the 5′-flanking sequence of CB1R identified 16 potential SF-1-bind-
In EMSAs, 13 of these 16 potential sites formed specific SF-1-related complexes that were supershifted by the addition of an anti-SF-1 antiserum (data not shown). The boxes in Fig. 4A depict the 13 sites in the CB1R 5-flanking region that bound SF-1.

We next performed transient transfection analyses to examine whether SF-1 could stimulate CB1R promoter activity. As shown in Fig. 4, B and C, SF-1 stimulated activity of CB1R sequences spanning −1565 to +48 (−1565/+48) in both Y1 cells and 293 cells. These data suggest that SF-1 can directly stimulate CB1R promoter activity.

To identify which, if any, of the 13 putative SF-1-responsive elements might mediate this induction, we made 5′-deletion constructs that included CB1R sequences from −1466 to +48 (−1466/+48), −1259/+48, −906/+48, −395/+48, −266/+48, −195/+48, and −75/+48 (Fig. 4A). We then analyzed promoter activities of these constructs via transient transfection in 293 cells (Fig. 4C). Deletion of 5′-flanking se-
Fig. 3. CNS Expression of SF-1 Contributes to CB1R-Dependent Effects on Nocturnal Food Intake and Body Weight
Effects of MA (A), ACEA (B), and AM251 (C) on nocturnal food intake in WT (SF-1\textsuperscript{flox/flox}\textsuperscript{Cre}\textsuperscript{+}+/+) and CNS-specific SF-1 KO (SF-1\textsuperscript{flox/flox}\textsuperscript{Cre}+/-) mice. After 24 h of fasting, mice were injected with either vehicle (veh), the CB1R agonists MA (1 mg/kg) or ACEA (3 mg/kg), or the CB1R antagonist AM251 (5 mg/kg), and nocturnal food intake was measured hourly thereafter for 3 h as described in Materials and Methods. Injection of MA or ACEA significantly stimulated nocturnal food intake in WT mice (*, P < 0.01, t test); their effects on appetite were insignificant in CNS-specific SF-1 KO (**, P > 0.1, t test). Treatment with the CB1R antagonist AM251 significantly inhibited food intake in both WT and SF-1 KO mice (*, P < 0.01, t test), but the effect was significantly greater in WT mice (‡, P < 0.05, t test). Graphs represent means ± SEM and data were analyzed by multifactorial ANOVA with repeated measure (strain × drug × time, P < 0.05).

Fig. 4. SF-1 Directly Regulates CB1R Expression
A, Schematic diagram of the 5'-flanking region of the mouse Cb1r gene, including the location of potential SF-1 binding sites. Shown below are the 5'-deletion constructs. B, CB1R promoter activity is significantly stimulated by SF-1 in mouse Y1 adrenocortical cells and human HEK 293 kidney cells in a dose-dependent manner (*, P < 0.001, t test). C, 5'-Deletion analysis of SF-1 binding sites that regulate the CB1R promoter. The 293 cells were cotransfected with SF-1 and the deletion constructs shown in Fig. 4A. Deletion of putative SF-1 binding sites in the -246 and -108 constructs significantly decreased promoter activity (**, P < 0.001, one-way ANOVA).
sequences from −1565 to −266 did not significantly affect SF-1 stimulated CB1R promoter activity. In contrast, deletion of sequences from −266 to −75 completely abolished SF-1 activation of promoter activity, implying that the sites at −233 and −101 may mediate SF-1 regulation (Fig. 4C).

To examine directly the roles of these two CB1R promoter sites in SF-1 regulation, we mutated the respective sequences in the context of the promoter region from −266/+48. Oligonucleotides that incorporated these mutations failed to bind SF-1 in gel mobility shift assays (Fig. 5A), whereas oligonucleotides containing native sequences at −233 and −101 specifically inhibited the SF-1 complex in a dose-dependent manner (Fig. 5A).

To further define the roles of the sites at −233 and −101 in SF-1 regulation of CB1R promoter activity, we examined the promoter activity and regulation by SF-1 using various mutated constructs (Fig. 5B). Mutation of the site at −233 alone did not significantly impair SF-1 induction of CB1R promoter activity, whereas mutation of both elements virtually abolished the response to SF-1. These results suggest that SF-1 directly stimulates CB1R promoter activity, predominantly via the element at −101 (Fig. 5C).

**SF-1 Regulates CB1R-Mediated ERK Activation Both in Cultured Cells and in Vivo**

CB1R is a G protein-coupled receptor that signals via G\(\alpha_{i/o}\) to activate cellular responses, including c-Jun N-terminal kinase (JNK), signal transducer and activator of transcription (STAT) 3, and the MAPK pathway (22–24). If CB1R is a bona fide target of SF-1, it is plausible that modulation of SF-1 could increase its expression in a manner that affected one of these CB1R-mediated signaling pathways. To address SF-1 regulation of CB1R-mediated cellular responses, we used mouse neuroblastoma Neuro2A cells, which endogenously express CB1R but do not express SF-1. First, we used real-time PCR assays to examine whether transfection of Neuro2A cells with SF-1 induces their expression of CB1R transcripts. These studies showed that introduction of SF-1 in the Neuro2A cells increased CB1R transcripts by approximately 4-fold (Fig. 6A).

Treatment of Neuro2A cells with MA (1 \(\mu M\)) increased phosphorylation of ERK, and this increase was significantly potentiated by SF-1 transfection (Fig. 6, B and C). Unlike the dynamic change of ERK phosphorylation by MA treatment, the levels of p-JNK and p-STAT3 were not affected by MA treatment (Fig. 6B). To determine whether G\(\alpha_{i/o}\) mediates the potentiation of ERK phosphorylation by SF-1 overexpression, we pretreated cells with pertussis toxin, which inhibits G\(\alpha_{i/o}\)-mediated signaling, or AM251, a CB1R antagonist. The potentiation of ERK phosphorylation by SF-1 overexpression was completely abolished by pretreatment with either pertussis toxin or AM251 (data not shown), indicating that the potentiation of ERK activation by SF-1 overexpression is CB1R agonist-dependent and requires the G\(\alpha_{i/o}\)-mediated pathway.

To further ascertain that the potentiation of ERK activation by SF-1 is directly mediated by CB1R in the Neuro2A cells, we showed that this effect was abolished after introduction of multiple siRNAs that specifically targeted CB1R (Kim, K., unpublished observation). These data collectively establish that pERK potentiation by SF-1 is directly mediated by CB1R.

Based on the potentiation of CB1R-mediated ERK phosphorylation by SF-1 in cultured cells, we wondered whether a similar effect also took place in vivo. Indeed, systemic administration of MA to WT (SF\(^+/\)) mice significantly increased p-ERK immunoreactivity in the VMH. Consistent with the ISH studies (Fig. 1, B and C), MA-induced phosphorylation of ERK in the VMH was markedly impaired in CNS-specific SF-1 KO mice (Fig. 7).

**DISCUSSION**

Results presented here show that SF-1 directly regulates CB1R via its 5’-flanking region and that the CNS-specific KO of SF-1 has significant effects on the response of food intake to exogenously administered CB1R agonist and antagonists. Specifically, we have demonstrated that CB1R expression was impaired in the VMH when we selectively removed SF-1 in the CNS, implicating SF-1 in CB1R expression in the VMH. We further found that CB1R colocalized with SF-1 expression in VMH neurons and that cannabinoids decreased excitability of SF-1 neurons but had no effect on non-SF-1 neurons. Recent evidence suggests that cannabinoid and leptin signals are integrated in lateral hypothalamic neurons (25). A similar integration may also occur in SF-1 neurons in the VMH, whose action potential frequency is inhibited by CB1R agonists (Fig. 2) but increased by leptin (14). These results suggest that integration of cannabinoid and leptin signals may regulate feeding behavior in the VMH.

Although SF-1 clearly is a key factor in VMH development, its specific roles in VMH function remain incompletely understood. We recently identified the CRH (Crh) receptor 2 as a direct target of SF-1, providing a plausible molecular basis for the increased anxiety-like behavior seen in CNS-specific SF-1 KO mice (7). Although CB1R in the brain also has been implicated in the regulation of anxiety, the key regions for emotional regulation apparently are the amygdala, hippocampus, and cortex. These sites do not express SF-1, and thus it cannot regulate their expression of CB1R. However, impairment of CB1R expression by genetic ablation of SF-1 may also contribute to the increased anxiety-like behavior because the VMH is known to be a critical brain region for defensive responses to predators (26, 27). Moreover, the acute administration of the CB1 antagonist SR141716A in-
Fig. 5. The SF-1-Responsive Element at −101 Is Required for SF-1 Regulation of CB1R Promoter Activity

A, EMSAs with the potential SF-1 binding sites at −233 and −101 in the CB1R promoter. Oligonucleotides comprising the CB1R promoter elements at −233 (left panel) and −101 (right panel) formed specific complexes with SF-1, whereas probes with mutation of the SF-1 core motif (M) failed to form this complex. *, Reactions that included 1 μl of an anti-SF-1 antiserum. Competition analysis revealed that the complexes formed with the −233 and −101 elements were specific. Unlabeled oligonucleotide competitors at the indicated molar excesses (2, 5, 10, and 100×) specifically inhibited SF-1 binding to the −233 and −101 elements, with complete abrogation of binding at the 100× molar excess. Oligonucleotides with mutations in the SF-1 binding motif (see Materials and Methods) did not inhibit the SF-1 complex. M, Mutated probe.

B, Schematic diagram of promoter constructs examined by transient transfection analyses. Empty squares indicate putative SF-1 binding sites and filled squares represent sites that were mutated. C, The SF-1 binding sites at −101 is crucial for SF-1 regulation. The indicated CB1R promoter constructs were transfected with or without SF-1 in 293 cells (left panel) and without SF-1 in Y1 cells (right panel). Mutation of the −233 site alone did not significantly impair SF-1 induction, whereas mutation of both the −233 and the −101 sites abolished SF-1 induction in 293 cells and markedly impaired promoter activity in Y1 cells (*, *P < 0.01, one-way ANOVA).
Produced anxiety-like responses in certain behavioral tests (28), and global CB1R KO showed increased anxiety-like behavior (29, 30).

Both N-methyl-D-aspartate subunit (31) and BDNF (32) have been identified as direct targets of SF-1 in the VMH.

Fig. 6. SF-1 Stimulates CB1R Expression and CB1R-Mediated ERK Phosphorylation in Neuro2A Neuroblastoma Cells

A, Transfection with SF-1 stimulates endogenous expression of CB1R transcripts in Neuro2A cells. Neuro2A cells were transiently transfected with either pcDNA3.1 or with the same vector expressing SF-1, and levels of CB1R transcripts were determined by real-time PCR assays as described in Materials and Methods. B, Transfection of Neuro2A cells with an expression plasmid for SF-1 selectively stimulates CB1R-mediated induction of ERK phosphorylation in Neuro2A cells. Immunoblot analyses showed that transfection of Neuro2A cells with an SF-1 expression plasmid dramatically induced MA-induced ERK phosphorylation at 5 and 10 min without affecting the total level of immunodetectable ERK, p-JNK, or p-STAT3.

Fig. 7. Administration of MA to WT But Not SF-1 KO Mice Significantly Stimulates ERK Phosphorylation in the VMH

WT and SF-1 KO mice were treated with MA and cells that contained phospho-ERK were quantified in sections of the mediobasal hypothalamus as residing in the dorsomedial (dm), central (c), or lateral (lat) regions of the VMH. A, Representative sections of immunoreactive phospho-ERK in vehicle- and MA-treated WT and CNS-specific SF-1 KO mice. B, The ERK phosphorylation induced by MA injection was significantly diminished in CNS-specific SF-1 KO mice. Bars represent means ± SEM (*, P < 0.01; **, P < 0.05, n = 3, t test).

Both N-methyl-D-aspartate subunit (31) and BDNF (32) have been identified as direct targets of SF-1 in the VMH.
A recent report expanded considerably the list of SF-1 targets in the VMH, including several cell adhesion molecules such as Amigo2, Cadh4, Sema3a, Slit3, and Netrin3 and other genes that are highly expressed in the VMH, such as Fez1, Npx2, Nkx2-2, and A2bp1 (11). Our results show that SF-1 can regulate CB1R expression directly, expanding the group of target genes to this G protein-coupled receptor, which has become an exciting target for drug development (33).

Given that CB1R activation by cannabinoids leads to phosphorylation of STAT3, JNK, and ERK (22, 23, 34), we sought to determine whether SF-1 has a role in signaling mediated by CB1R activation. As shown in Fig. 6, transfection of Neuro2A cells with SF-1 induced CB1R expression and potentiated CB1R-mediated ERK phosphorylation. Activation of STAT3 or JNK by CB1R has been implicated in neurite outgrowth (22–24), but the precise effects of CB1R-dependent ERK phosphorylation after CB1R activation are not fully understood. Activation of ERK has been related to the regulation of proliferation and differentiation (35), and this pathway, like SF-1 itself, may play important roles in the differentiation/migration of VMH neurons. Studies presented here show that systemic administration of a CB1R agonist is associated with increased levels of immunoreactive phospho-ERK in VMH neurons, suggesting that CB1R also couples to the MAPK pathway in the VMH (Fig. 7).

Although behaviors modulated by cannabinoids are regulated by complex interactions among structures such as the ventral tegmental area, medial forebrain bundle, nucleus accumbens, and amygdala (36), the hypothalamic nuclei surrounding the third ventricle are central to CB1R-mediated energy balance (37–39). For example, direct administration of the CB1 agonist anandamide (ANA) into the VMH significantly increased food intake, whereas the CB1R antagonist SR141716 inhibited the AEA induction of food intake (3) and infusion of ANA into the nucleus accumbens showed hyperphagic effects with marked activation of hypothalamic nuclei involved in feeding behaviors (39). These results implicate hypothalamic CB1R in orexigenic actions of cannabinoids. In agreement with these studies, we observed that administration of CB1R agonists (e.g. MA and ACEA) increased food intake in WT mice (Fig. 3, A and B); this increased food intake was blunted significantly in mice with CNS-specific KO of SF-1 (Fig. 3, A and B). Likewise, administration of the specific CB1R antagonist AM251 markedly suppressed food intake in WT mice, but suppression was significantly impaired in CNS-specific SF-1 KO mice (Fig. 3C). These data imply that the lack of CB1R expression in VMH neurons may be a factor in the behavioral effects exhibited by CNS-specific SF-1 KO mice (e.g. increased anxiety-like behavior and adiposity) (6, 7).

A number of reported interactions between the cannabinoid system and neuropeptides involved in energy homeostasis provide insights into potential roles of the cannabinoid system on food intake and hypothalamic regulation. For example, SF-1 neurons in the VMH express leptin receptor and leptin signaling in SF-1 neurons is required for normal energy homeostasis (13, 14); similarly, these SF-1 neurons express CB1R and SF-1 directly regulates the CB1R promoter (this study). Leptin negatively regulates endocannabinoid production (37) and modulates food intake by inhibiting the orexigenic neuropeptide Y (NPY) in arcuate nucleus (40, 41). There are conflicting data regarding the cannabinoid system and neuropeptide NPY regulation. Stimulation of the cannabinoid system with ANA and CP55,940 reportedly augmented NPY release in the rat hypothalamus (42), suggesting that activation of the cannabinoid system in the hypothalamus generated orexigenic effects by augmenting NPY levels. In contrast, others have proposed that cannabinoids modulate energy balance by NPY-independent mechanisms (37). Consistent with the latter model, hypothalamic NPY expression was not affected by activation of the cannabinoid system by the CB1R agonist MA (data not shown) or by the perturbed VMH expression of CB1R in the VMH in CNS-specific SF-1 KO mice. Rather, the orexigenic effect of cannabinoids may involve other neuropeptides such as CRH (43), cocaine-amphetamine regulated peptide (44), proopiomelanocortin (45), and ghrelin (46).

In summary, our data identify CB1R as a novel target of SF-1 in the VMH, providing clear evidence that SF-1 plays important roles in the regulation of CB1R expression as well as in CB1R-mediated regulation of food intake. Furthermore, the excitability of SF-1 neurons is directly inhibited by CB1R agonists, which may be an important factor in homeostatic regulation by the VMH. Future studies that identify downstream modulators whose activity is controlled by cannabinoid signaling in SF-1 neurons will provide new insights into how SF-1 maintains CB1R-mediated energy balance in the VMH and will hopefully identify other genes that determine how the VMH and other hypothalamic nuclei regulate various physiological functions.

MATERIALS AND METHODS

Animal Care

All mouse care and experimental procedures were approved by the Institutional Animal Care Research Advisory Committee at UT Southwestern or at Albert Einstein Medical Center. Mice were housed at room temperature with a 12-h light, 12-h dark cycle (lights on at 0600 h) with regular mouse chow (Teklad mouse/rat diet 7001; 3.82 kcal/g, gross energy, 4.25% kcal from fat) and water provided ad libitum. The generation of CNS-specific SF-1 KO (SF-1^FKO; F^+^−) mice was described previously (7).

Drugs

The CB1 agonists, MA and ACEA, and the CB1 antagonist AM251 were purchased from Sigma-Aldrich (St. Louis, MO). MA was dissolved in saline solution containing 2% polyoxyethylene sorbitan monolaurate (Tween 80; Sigma) and 20% ethanol. ACEA was suspended in saline solution with 1% Tween 80, and AM251 was dissolved in saline containing 2% Tween 80 and 10% dimethylsulfoxide. All drugs were injected ip in a volume of 1 µg/g body weight.
Nocturnal Food Intake and Body Weight Studies

The WT (SF-1<sup>f<sup>Cre<sup>/</sup></sup>), heterozygous (SF-1<sup>f<sup>/+/</sup></sup>), and CNS-specific SF-1 KO (SF-1<sup>f<sup>Centre</sup>/</sup>) mice were individually housed beginning at 4 wk of age. Body weight and food intake were monitored three or four times per week until 9 wk of age. Matched littersmates with comparable food intake and weights (21–23 g for males and 18–19 g for females) were used for these studies. At d 65, food was removed from the cages at 1800 h and mice were fasted for 24 h; thereafter, they were administered the indicated drugs via ip injection and food intake was monitored over 3 h.

Slice Preparation for Electrophysiology

Transverse brain slices were prepared from SF-1/eGFp mice at postnatal age 21-28 d. Animals were anesthetized with a mixture of ketamine and xylazine. After decapsulation, the brains were transferred into a sucrose-based solution bubbled with 95% O<sub>2</sub>-5% CO<sub>2</sub> and maintained at approximately 3 C. This solution contained (mM): sucrose 248; KCl2; MgCl<sub>2</sub> 1; KH<sub>2</sub>PO<sub>4</sub> 1.25; NaHCO<sub>3</sub> 26; sodium pyruvate 1; and glucose 10. Transverse coronal brain slices (200 μm) were prepared using a vibratome. Slices were equilibrated with an oxygenated artificial cerebrospinal fluid (aCSF) for more than 1 h at 32 C before transfer to the recording chamber. The slices were continuously superfused with aCSF at a rate of 1.5 ml/min containing (in mM): NaCl 113; KCl<sub>3</sub>; NaH<sub>2</sub>PO<sub>4</sub> 1; NaHCO<sub>3</sub> 26; CaCl<sub>2</sub> 2.5; MgCl<sub>2</sub>; and glucose 5 in 95% O<sub>2</sub>-5% CO<sub>2</sub> at room temperature.

Electrophysiological Studies

Brain slices were placed on the stage of an upright, infrared-differential interference contrast microscope (Olympus BX50WI, Olympus America Inc., Center Valley, PA) mounted on a Gibraltar X-Y table (Burleigh, Olympus America Inc.) and visualized with a ×40 water immersion objective by infrared microscopy (DAGE MTF camera; DAGE, Michigan City, IN). Membrane potentials were recorded at room temperature (25–26 C) with an Axopatch 200B Patch-Clamp amplifier. CNQX (10 μM), DL-amino-phosphononvaleric acid (DL-AP-5, 50 μM), picrotoxin (100 μM) and strychnine (1 μM) were continuously present in aCSF. The internal solution contained (mM): K acetate 115; KCl 10; MgCl<sub>2</sub> 2; EGTA 0.2; HEPES 10; Na<sub>2</sub>ATP 1; Na<sub>2</sub>GTP 0.5; and phosphocreatine 5. Pipette resistance ranged from 2.5 to 4 MΩ. WIN 55212-2 and AM251 were purchased from Sigma-Aldrich and dissolved in dimethylsulfoxide at 10,000× the final concentration. Effects of the CB1R agonist compared with control were analyzed using Student’s t tests (Origin 7.0). Differences were considered significant when the P value was < 0.05. All statistical results are given as means ± SEM.

Immunohistochemistry and ISH

Immunohistochemistry and ISH were performed as described (7). Briefly, floating or mounted sections from adult brain (age 12–18 wk if not specified) were used for immunohistochemistry and ISH. Images were captured with a Nikon E1000 automated microscope connected with a Nikon digital camera (DXM1200F, Nikon, Melville, NY). Rabbit anti-SF-1 antibody [1:1500, (47)] was used for SF-1 immunohistochemistry. Samples were incubated with a biotin-conjugated antirabbit secondary antibody (1:1000) from Vector Laboratory (Burlingame, CA; BA-1000) for 2 h, followed by incubation for 1 h in a solution of avidin-biotin complex (Vectastain Elite ABC kit, Vector Laboratory). Finally, signals were visualized with DAB substrate kit (SK-4100; Vector Laboratory).

For immunofluorescent analysis of pERK activation in the VMH, mice at 9 wk were fasted for 24 h and then fed for 1 h after MA (1 mg/kg) administration. Mice were anesthetized with tribromoethanol (Avertin, 250 mg/kg-BW) and perfused via cardiac puncture with 4% paraformaldehyde (Sigma) in PBS (pH 7.4). After cryoprotection using 30% sucrose, brains were sectioned at 20 μm on a cryostat (Leica, Bannockburn, IL; CM1900). Rabbit anti-phosphorylated (p)-ERK antibody (1:100, no. 4376; Cell Signaling Technology, Danvers, MA) was incubated overnight and followed by Cy3-conjugated goat anti-rabbit IgG (Jackson ImmunoResearch, West Grove, PA) as secondary antibody.

ISH for CB1R was performed on every fourth serial section from at least five brains. The mouse CB1R probe was generously provided by Dr. Joel Elmquist (UT-Southwestern, Dallas, TX) and included CB1R sequences from 953 to 1557.

Cell Culture and Transfection Analyses

Cell culture conditions were previously described (7). HEK 293 human embryonic kidney cells were maintained in DMEM supplemented with 10% fetal bovine serum (FBS) and 100 U/ml penicillin-streptomycin (Invitrogen, Carlsbad, CA). Neuro2A mouse neuroblastoma cells were cultured using MEM (Invitrogen) containing nonessential amino acids, 10% FBS, and 100 U/ml penicillin-streptomycin. Y1 mouse adrenocortical tumor cells were grown in Ham's F10 medium (Mediatech, Herdon, VA) supplemented with 15% horse serum, 5% FBS, and 100 U/ml penicillin-streptomycin.

For immunoblotting analyses, Neuro2A cells were plated in six-well culture plates at a density of 5 × 10<sup>5</sup> cells per well and cultured in the growth media for 20–24 h before transfection. Cells were transfected with Fugene 6 (Roche Diagnostics, Indianapolis, IN) according to the manufacturer’s instructions. After 24 h, cells were treated with MA for differing time periods. Cells were then lysed in 100 μl of lysis buffer containing protease and phosphatase cocktails (Roche Diagnostics) and followed by gentle shaking on ice for 10 min. Supernatants were collected and resolved on 10–12% sodium dodecyl sulfate-polyacrylamide gels for 2 h and then transferred to nitrocellulose membrane (Bio-Rad, Hercules, CA). Commercially available antibodies from Cell Signaling Technology were used; rabbit anti-p-ERK (1:1000; no. 4376), rabbit anti-ERK (1:2000; no. 9102), rabbit anti-pJNK (1:2500; no. 9251), and rabbit anti-pS9 (1:1000; no. 9131) for primary reaction. Signals were detected by horseradish peroxidase-conjugated secondary antibodies. The chemiluminescence was visualized with Lumi-Light Western Blotting Substrate (Roche Diagnostics), and the images were detected by Kodak Biomax MR film. The bands were quantified with the ImageJ program (National Institutes of Health; http://rsb.info.nih.gov/ij/).

EMSA

Duplex oligonucleotides containing the putative SF-1 binding sites at −233 and −101 were labeled with <sup>32</sup>P by fill-in with Klenow fragment (New England Biolas, Newton, MA) and used as probes for EMAS as described (7). Oligonucleotide probes were 5'-GGGGCGGCAACCGAGCCGCTCCCCAGCTAGGGAGGGACGGGCGCCGTCGCCTTCATG-3' for the site −233 (WT, top strand), 5'-GCTAGGAGGAGCGCCTTGCGCTGCAGCCACT-3' for the site −233 (WT, bottom strand), 5'-GGGCCGAGGCCCAAAACCCCTCCCTCAGTTTAG-3' for site −233 (M, top strand), 5'-GCTAGGAGGAGCGCCTTGCGCTGCAGCCACT-3' for the site −233 (M, bottom strand), 5'-GGGGCGGCAACCGAGCCGCTCCCCAGCTAGGGAGGGACGGGCGCCGTCGCCTTCATG-3' for the site −101 (WT, top strand), 5'-GAATGAGCCTTTGCTCTGCGTCCAGCGAAGGAGGCAAAGGTCCTATT-3' for site −101 (WT, bottom strand), 5'-GCTAGGAGGAGCGCCTTGCGCTGCAGCCACT-3' for site −101 (M, top strand), 5'-GAATGAGCCTTTGCTCTGCGTCCAGCGAAGGAGGCAAAGGTCCTATT-3' for site −101 (M, bottom strand). Boldface represents SF-1 binding sites and mutated sequences. Unlabeled duplex oligonucleotides were used for competition experiment at 2-, 5-, 10-,
and 100-fold molar excesses. Where indicated, a rabbit antitoxin SF-1 antisera (1 μl) was included in the binding reaction. Nuclear extract was prepared from mouse Y1 adrenocortical tumor cells, which endogenously express SF-1, as described (7).

Real-Time Q-PCR Analysis of RNA from Purified Hypothalamic Cells Expressing a SF-1/eGFP Transgene

Embryonic d 16.5 embryos were harvested and cells were prepared as described (12). Total RNA was prepared from 10,000–15,000 fluorescence-activated cell sorting-enriched cells using TRIzol (Invitrogen), deoxyribonuclease I treatment (TAAGEN, Valencia, CA), and RNeasy columns (TAAGEN). 2.5–25 ng of RNA was then amplified using the WT-Pico Kit (NuGEN, San Carlos, CA). In a final volume of 10 μl, the real-time PCR assays contained 1 ng of amplified RNA, 150 nM of each primer, and 5 μl of SYBR Green PCR Master Mix (Applied Biosystems, Foster City, CA). PCRs were performed on an Applied Biosystems 7900HT and relative mRNA levels were calculated using the comparative threshold cycle method (Applied Biosystems user bulletin no. 2). Comparisons were made to samples from cells harvested from WT embryos using cyclophilin as the reference gene. Real-time primers were designed using Primer Express software (PerkinElmer Life Sciences, Downers Grove, IL) and validated using a standard curve standard to cyclophilin (see supplemental figure published as supplemental data on The Endocrine Society’s Journals Online web site at http://mend.endojournals.org). Primers used were forward (F) and reverse (R) CB1R (F) 5′-GTGCCGAGGAGCTTCTG-3′ and (R) 5′-TCTTTGTAGGAGCCAGCTCACA-3′, Cyclophilin (F) 5′-TG-GAGAAGCACCAGACAGACA-3′ and (R) 5′-TGCGGAAGTCCACAAATG-3′. Twelve different pools (three WT male, three KO male, three WT female, and three KO female) were analyzed. Because no sexual dimorphism was noted, the data were combined as WT (n = 6) or KO (n = 6).

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