Hypocretin-2-Saporin Lesions of the Lateral Hypothalamus Produce Narcolepsy-Like Sleep Behavior in the Rat

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Hypocretins (Hcrts) are recently discovered peptides linked to the human sleep disorder narcolepsy. Humans with narcolepsy have decreased numbers of Hcrt neurons and Hcrt-null mice also have narcoleptic symptoms. Hcrt neurons are located only in the lateral hypothalamus (LH) but neither electrolytic nor pharmacological lesions of this or any other brain region have produced narcoleptic-like sleep, suggesting that specific neurons need to be destroyed. Hcrt neurons express the Hcrt receptor, and to facilitate lesioning these neurons, the endogenous ligand hypocretin-2/orexin B (Hcrt2) was conjugated to the ribosome-inactivating protein saporin (SAP). In vitro binding studies indicated specificity of the Hcrt2-SAP because it preferentially bound to Chinese hamster ovary cells containing the Hcrt/orexin receptor 2 (Hcrtr2/OX2R) or the Hcrt/orexin receptor 1 (Hcrtr1/OX1R) but not to Kirsten murine sarcoma virus transformed rat kidney epithelial (KNRK) cells stably transfected with the substance P (neurokinin-1) receptor. Administration of the toxin to the LH, in which the receptor is known to be present, eliminated some neurons (Hcrt, melanin-concentrating hormone, and adenosine deaminase-containing neurons) but not others (a-melanocyte-stimulating hormone), indicating specificity of the toxin in vivo. When the toxin was administered to the LH, rats had increased slow-wave sleep, rapid-eye movement (REM) sleep, and sleep-onset REM sleep periods. These behavioral changes were negatively correlated with the loss of Hcrt-containing neurons but not with the loss of adenosine deaminase-immunoreactive neurons. These findings indicate that damage to the LH that also causes a substantial loss of Hcrt neurons is likely to produce the multiple sleep disturbances that occur in narcolepsy.

Key words: hypothalamus; peptides; lesion; sleep; REM sleep; circadian rhythm

Regular periods of sleep and wakefulness occur in virtually all mammals and birds. Sleep is generally divided into two states, slow-wave sleep (SWS) and rapid-eye movement (REM) sleep. The neuronal mechanisms underlying the regulation of sleep and wakefulness are unclear (for review, see Shiromani, 1998), and to gain a better understanding investigators have studied the human sleep disorder narcolepsy. Narcolepsy is a disabling neurological disorder characterized by excessive daytime sleepiness, sleep attacks, sleep fragmentation, cataplexy, sleep-onset REM sleep periods (SOREMP), and hypnagogic hallucinations. Recently, narcolepsy was linked with the loss of neurons containing the hypocretin (Hcrt) neuropeptides (Peyron et al., 2000; Thannickal et al., 2000). Narcoleptic patients have low CSF concentrations of hypocretin-1 (Nishino et al., 2000), which is consistent with a decrease in the number of Hcrt neurons. Moreover, narcoleptic canines possess a mutation in the hypocretin-2 receptor (Lin et al., 1999), and mice with deletion of the Hcrt gene exhibit symptoms of narcolepsy (Chemelli et al., 1999).

The hypocretin peptides, also known as the orexins, are produced exclusively by neurons located in the lateral hypothalamus (LH) (De Lecea et al., 1998; Peyron et al., 1998; Sakurai et al., 1998). A single gene encodes Hcrt, which is cleaved by proteolytic processing into two smaller peptides, hypocretin-1 (orexin A) and hypocretin-2 (orexin B) (De Lecea et al., 1998; Sakurai et al., 1998). Hypocretin/orexin-containing neurons project to the entire brain and spinal cord (De Lecea et al., 1998; Date et al., 1999; Nambu et al., 1999).

Hcrt-containing neurons are located in a part of the brain that von Economo considered to be a “wake” center (von Economo, 1930). However, his findings were ignored because lesions of the posterior hypothalamus have to date yielded inconsistent effects on sleep and wakefulness. For instance, Ranson (1939), Nauta (1946), and Shoham and Teitelbaum (1982) observed behavioral signs of sleepiness after electrolytic lesions of the posterior hypothalamus but did not report the daily amounts of sleep, which makes it difficult to conclude whether the behavioral symptoms were isolated or pervasive. When long-term electroencephalogram (EEG) sleep recordings were made, increased wakefulness and reduced REM sleep were obtained after electrolytic lesions of the LH (McGinty, 1969; Dangui and Nicolaids, 1980; Jurkowitz et al., 1994). Swett and Hobson (1968) reported increased SWS accompanied by behavioral rigidity in some cats, but their electrolytic lesions encompassed the ventral tegmental area. Excitotoxic lesions of the posterior hypothalamus with ibotenic acid have also produced transient effects, such as increased sleep for 1–4 d followed by increased wakefulness; REM sleep was...
increased only during the first 3–21 hr (Sallanon et al., 1988; Denoyer et al., 1991).

The inconsistent effects on sleep after electrolytic or excitotoxic lesions of the posterior hypothalamus or LH might have occurred because the methods used to make the lesion did not destroy the appropriate neurons. Moreover, some neurons are resistant to lesion by excitotoxins such as ibotenate (Yanai et al., 1997). In the present study, to more effectively target the Hcrt system, the ribosome-inactivating protein saporin (SAP) (Stirpe et al., 1992) was conjugated to the hypocretin/orexin receptor binding ligand hypocretin-2/orexin-B (Hcrt2) to lesion Hcrt receptor-bearing neurons. The LH contains a high concentration of Hcrt receptor mRNA (Trivedi et al., 1998) and immunoreactivity (Hervieu et al., 2001), and the Hcrt-immunoreactive (ir) axons make synaptic contacts with Hcrt-containing perikarya (Hovrath et al., 1999), indicating the presence of the Hcrt receptor on Hcrt neurons. When the Hcrt2-SAP was administered to the LH of rats, the toxin lesioned LH neurons, including the Hcrt-ir neurons, and produced symptoms that are characteristic of narcolepsy. These findings identify the LH as promoting wakefulness and inhibiting REM sleep and demonstrate that the Hcrt2-SAP conjugate is a useful tool for investigating the Hcrt system.

MATERIALS AND METHODS

Experiment 1: In vitro analysis of binding of Hcrt2-SAP

Hypocretin receptor-containing cells. Stably transfected cell lines expressing the hypocretin/orexin receptor 1 (HcrtR1/OX1R) or receptor 2 (HcrtR2/OX2R) are described by Sakurai et al. (1998). Chinese hamster ovary cells expressing HcrtR1/OX1R or HcrtR2/OX2R (gifts from Dr. M. Yanagisawa, University of Texas Southwestern Medical Center, Dallas, TX) were cultured in DMEM supplemented with 10% fetal calf serum at SRI International. For fixation, 2.5 × 10^6 cells/sample were washed with 1 ml of fluorescent-activated cell sorting (FACS) buffer (2% fetal bovine serum in PBS) per 10^6 cells. The buffer was then removed and cells were fixed by resuspension of the pellet in 1% paraformaldehyde (1 ml/10^6 cells). After a 15 min incubation at 4°C, an equal volume of FACS buffer was added; cells were pelleted after thorough mixing. The pellets were washed as described above and resuspended in ice-cold 90% ethanol (1 ml/10^6 cells). After a 1 hr incubation at 4°C, an equal volume of FACS buffer was added; cells were pelleted after mixing. The pellets containing the fixed cells were washed again, resuspended in 200 μl of FACS buffer per sample (2.5 × 10^6 cells), and shipped to Advanced Targeting Systems for analyses.

To identify whether the Hcrt2-SAP bound to another peptide receptor, Kirsten murine sarcoma virus transformed rat kidney epithelial (KNRK) cells stably transfected with the substance P [neurokinin-1 (NK-1)] receptor (a gift from Dr. Nigel Bunnett, University of California, San Francisco, CA) were used (Wiley and Lappi, 1997).

FACS analysis. FACS analysis was performed at Cytometry Research LLC (San Diego, CA). Adherent cells were detached using CellStripper (Cellgro; Mediatech, Herndon, VA) and counted. A total of 2.5 × 10^6 cells/sample were washed with FACS buffer. Hcrt2-SAP or substance P attached to saporin (SP-SAP) was applied to the cells at a final concentration of 100 nM in 200 μl of FACS buffer. Samples were incubated for 1 hr at 4°C. Samples were washed twice with 1 ml of FACS buffer. A chicken anti-saporin antibody (Advanced Targeting Systems) was applied at a dilution of 1:50 in 100 μl of FACS buffer. Samples were incubated for 1 hr at 4°C and then washed as described previously. Rabbit anti-chicken IgY conjugated to FITC (Chemicon, Temecula, CA) was applied at a 1:50 final dilution in 100 μl of FACS buffer. Samples were incubated for 30 min at 4°C and then washed as described previously. Cells were resuspended in 500 μl of FACS buffer and then run on a FACS can (Becton Dickinson, Richmond, CA). Data were analyzed using CellQuest software (Becton Dickinson).

Experiment 2: Time course of the effects of Hcrt2-SAP on hypothalamic neurons

Subjects. Male Sprague Dawley rats (400–450 gm) (Charles River Laboratories, Wilmington, MA) were housed singly in Plexiglas cages with wood shavings; food and water were available ad libitum. The temperature in the room was 25°C and a 12 hr light/dark cycle (lights on from 7 A.M. to 7 P.M.; 100 lux) was maintained.

In 10 rats (under anesthesia with a cocktail of 0.75 mg/kg acpomazine, 2.5 mg/kg xylazine, and 22 mg/kg ketamine, i.m.), a unilateral injection of Hcrt2-SAP was made to the LH using a stereotaxic instrument; the rats were killed 2 (n = 3), 4 (n = 4), or 12 (n = 3) days later. The rats were perfused (after an overdose of Nembutal) with saline (100 ml) followed by 10% formalin (350 ml). The brains were carefully removed, placed overnight in the formalin solution, and then equilibrated in 30% sucrose solution at 4°C.

Microinjection sites. The Hcrt2-SAP conjugate (490 ng/0.5 μl; Advanced Targeting Systems) or pyrogen-free saline were delivered (0.5 μl) (Picospritzer; General Valve, Fairfield, NJ) using a glass micropipette (tip diameter of 20 mm). After injection the pipette was left in place for 5 min and then withdrawn slowly. A single injection was made in each rat in the LH (coordinates relative to bregma: anterior, −3.3 to −3.8 mm; lateral, 1.3–1.6 mm; ventral, 8.2–9.0 mm below the dura).

Immunohistochemistry. Tissue sections (30 mm thick) cut on a sliding microtome were incubated overnight at room temperature in the primary antibody (a one in five series for each primary antibody). After washing, the sections were placed in secondary and tertiary antibody (1:250) (Chemicon) followed by incubation in avidin–biotin complex for 1 hr (Vector Laboratories, Burlingame, CA). The DAB method was used to visualize the reaction product. The tissue sections were then counterstained with a Nissl stain (Neutral Red), dehydrated in graded alcohols, and coverslipped. Control sections were reacted without the primary antibodies or in preabsorbed serum; no labeled neurons were evident. The specificity of the antibodies was further confirmed by the restriction of the labeled neurons to the posterior hypothalamus.

Antibodies. Rabbit anti-orexin-A (hypocretin-ir) (1:70,000; Amersharm Pharmaica Biotech, Arlington Heights, IL), rabbit anti-adenosine deaminase (1:10,000; Chemicon); rabbit anti-melanin-concentrating hormone (MCH) (1:50,000; Chemicon), and rabbit anti-n-melanocyte-stimulating hormone (a-MSH) (1:5000; Chemicon) were purchased. Serial sections were incubated for 1 hr with the primary antibody (1:100,000; Chemicon) followed by incubation in avidin–biotin complex for 1 hr (Vector Laboratories, Burlingame, CA). The DAB method was used to visualize the reaction product. The tissue sections were then counterstained with a Nissl stain (Neutral Red), dehydrated in graded alcohols, and coverslipped. Control sections were reacted without the primary antibodies or in preabsorbed serum; no labeled neurons were evident. The specificity of the antibodies was further confirmed by the restriction of the labeled neurons to the posterior hypothalamus.

Experiment 3: Effects of bilateral injection of Hcrt2-SAP on sleep and wakefulness

Subjects. Twenty male Sprague Dawley rats were used in this experiment; housing conditions were the same as those described for experiment 2.

Surgery. The rats were implanted under anesthesia (cocktail of 0.75 mg/kg acpomazine, 2.5 mg/kg xylazine, and 22 mg/kg ketamine, i.m.) with screws (2.5 mm in diameter) to record EEG and electromyogram (EMG), as described previously (Shiromani et al., 2000). At least 2 weeks prior to the surgery, the rats were transilluminated, and, in each animal, the field drawings were made to identify the distribution of Hcrt, MCH, a-MSH, and ADA-ir neurons in the posterior hypothalamus using the Neurolucida program (Colchester, VT).

Analysis of sleep–wake states. Contralateral frontal–occipital EEG screw electrodes were used for EEG acquisition. The EEG data were filtered at 70 Hz (low-pass filter) and 0.3 Hz (high-pass filter) using a Grass electroencephalograph (Grass Instruments, Quincy, MA) and were continuously sampled at 128 Hz. The 24 hr EEG and EMG recordings were collected for at least 2 weeks. Next the rats were perfused (after overdose of Nembutal), and formalin-fixed brains were used for histological analysis.
activity. Slow-wave sleep consisted of high-amplitude slow waves together with a low EMG tone relative to waking. REM sleep was identified by the presence of desynchronized EEG and/or $\theta$ activity coupled with low EMG relative to slow-wave sleep. The amount of time spent in wakefulness, SWS, and REM sleep was determined for each hour. To determine whether there was a change in the amplitude of the diurnal activity. In young normal Sprague Dawley rats, the duration of wakefulness, SWS, and REM sleep was determined for each hour. To determine whether there was a change in the amplitude of the diurnal activity, SWS, and REM sleep was calculated using the ICLUS software system (M. Opp, University of Michigan, Ann Arbor, MI). After the EEG data were scored, the code was broken to reveal the identity of each rat. ANOVA and $t$ tests with Bonferroni corrections (where appropriate) were used to compare changes in sleep parameters.

The criteria used to identify SOREMPs were based on a combination of electrophysiological and behavioral observations and were modeled after those used in humans, because no such criteria exist for rats. In humans, SOREMPs are defined as episodes of REM sleep occurring within a 15 min window after the onset of sleep (Carskadon et al., 1986). In young normal Sprague Dawley rats, the duration of wake (day, 1.92 ± 0.1 min; night, 0.33 ± 0.47; data from Shiromani et al., 2000) and SWS (day, 3.83 ± 0.32; night, 2.8 ± 0.25; data from Shiromani et al., 2000) bouts are considerably shorter, making it necessary to modify the criteria. Accordingly, a SOREMP in the rat was identified as a REM sleep episode in the day or night that occurred after ≥2 min of wakefulness with <2 min of an intervening episode of SWS. The 2 min duration of wake and SWS bouts was based on the duration of these bouts in normal Sprague Dawley rats (Shiromani et al., 2000), and this duration was also observed in the saline-treated rats in the present study (Table 2). In addition to these electrophysiological criteria, a behavioral determination of a SOREMP was made when the videotape showed that the rat was lying down, had irregular respiration, and had phasic motor twitches. A Sony video camera (CCD-TRV16; Sony, Tokyo, Japan) with the capability to record in darkness was used to record the animal’s behavior. In the video clips, the EEG and EMG are superimposed on the behavior to facilitate identifying the behavioral state of the rat.

**Immunohistochemistry and cell counts.** The tissue was reacted for visualization of Hcrt or adenosine deaminase-ir neurons as described in experiment 2. The cell counts were performed as noted in experiment 2 by a person (M. Malik) who was blind to the type of drug administered. In this experiment, comparisons were made with the saline-injected rats.

### RESULTS

#### Experiment 1: In vitro analysis of binding of Hcrt2-SAP

FACS analysis was performed on cells transfected with the Hcrt receptors or NK-1 receptor. In this assay, the fluorescent probe was attached to antibodies to saporin, such that the entire Hcrt2-SAP was bound to substance P receptor (NK-1)-containing KNRK cells. In this assay, the fluorescent probe was attached to antibodies to saporin, such that the entire Hcrt2-SAP was bound to substance P receptor (NK-1)-containing KNRK cells. As expected, SP-SAP bound to a lesser degree, to the HcrtR1/OX1R (Fig. 1B). These data demonstrate that Hcrt2-SAP complex must be intact for a read-out to occur. Figure 1 shows that the Hcrt2-SAP binds to the HcrtR2/OX2 receptor and, to a lesser degree, to the HcrtR1/OX1 (Fig. 1A). These data are consistent with the properties of the ligand alone (Sakurai et al., 1998). There is no binding of Hcrt2-SAP to cells that are transfected with the substance P receptor, indicating lack of cross-reactivity with another peptide receptor. As expected, SP-SAP bound to substance P receptor (NK-1)-containing KNRK cells (Manthy et al., 1997) (Fig. 1B). These data demonstrate that binding of Hcrt2-SAP is specific for the Hcrt receptors.

### Table 1. Changes in light–dark ratios in rats with lateral hypothalamic injections of saline or Hcrt2-SAP

<table>
<thead>
<tr>
<th>Group</th>
<th>6 d after injection</th>
<th>14 d after injection</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>Wake</td>
<td>SWS</td>
</tr>
<tr>
<td>Hcrt cell loss &gt;60% ($n = 9$)</td>
<td>1.00 ± 0.12*</td>
<td>1.25 ± 0.15*</td>
</tr>
<tr>
<td>Hcrt cell loss &lt;30% ($n = 3$)</td>
<td>0.38 ± 0.09</td>
<td>3.56 ± 1.00</td>
</tr>
<tr>
<td>Saline ($n = 8$)</td>
<td>0.43 ± 0.03</td>
<td>2.99 ± 0.28</td>
</tr>
</tbody>
</table>

Values (±SEM) represent ratio of wakefulness, SWS, or REM sleep percentage during light-on period versus its corresponding value during light-off period. For EEG $\delta$ power, the values represent the ratio of $\delta$ power during the 2 hr period after lights on (7 A.M. to 9 A.M.) versus the 2 hr period after lights off (7 P.M. to 9 P.M.). For $\delta$ power, the 2 hr time points represented the peak and nadir of the $\delta$ power across the 24 hr in saline-treated rats, a finding that is consistent with the waxing and waning of EEG $\delta$ power (Borbely, 1994). *p < 0.05, significant difference compared with the values of the saline group.

### Table 2. Average (±SEM) number of transitions to SWS, REM sleep, or wakefulness and duration of wakefulness, SWS, or REM sleep in rats administered Hcrt2-SAP in the LH

<table>
<thead>
<tr>
<th>Group</th>
<th>Average number of transitions</th>
<th>Average duration of bouts (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>W-SWS</td>
<td>SWS-REM</td>
</tr>
<tr>
<td>Light-off period</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hcrt cell loss &gt;60% ($n = 9$)</td>
<td>93.7 ± 8.8*</td>
<td>41.2 ± 3.9</td>
</tr>
<tr>
<td>Hcrt cell loss &lt;30% ($n = 3$)</td>
<td>46.7 ± 8.8</td>
<td>24.7 ± 5.2</td>
</tr>
<tr>
<td>Saline ($n = 8$)</td>
<td>56.3 ± 4.6</td>
<td>30.6 ± 4.2</td>
</tr>
<tr>
<td>Light-on period</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hcrt cell loss &gt;60% ($n = 9$)</td>
<td>98.5 ± 7.1</td>
<td>20.5 ± 2.8*</td>
</tr>
<tr>
<td>Hcrt cell loss &lt;30% ($n = 3$)</td>
<td>97.0 ± 6.6</td>
<td>51.0 ± 6.4</td>
</tr>
<tr>
<td>Saline ($n = 8$)</td>
<td>116.0 ± 5.0</td>
<td>70.1 ± 3.6</td>
</tr>
</tbody>
</table>

W, Wakefulness. *p < 0.05, significance for each group compared with its respective saline group.
Experiment 2: Time course of the effects of Hcrt2-SAP on hypothalamic neurons

To identify the cytotoxic effects in vivo, the Hcrt2-SAP was administered unilaterally to the hypothalamic regions known to contain the Hcrt receptor. The neurons in the TMN were counted because they do not contain Hcrt but possess the receptor, as demonstrated by the presence of Hcrt receptor mRNA in the TMN (Fig. 2C). The TMN neurons represent a homogenous population of densely packed neurons that contain the enzyme ADA (Senba et al., 1985) (Fig. 2A, B), and Hcrt fibers densely innervate this nucleus (Fig. 2D).

Figure 3 summarizes the time-dependent loss of specific markers of neuronal phenotypes in the posterior hypothalamus. Comparisons made to the contralateral un.injected side indicated that rats killed on day 2 after injection (n = 3) had little loss of H-ir or ADA-ir neurons. However, by day 4 after injection there was a significant decrease (31%) in the number of Hcrt-ir neurons (paired t test with contralateral nonlesioned side, t = 6.07; df = 3; p < 0.009; power = 96.8). By day 12 there was a 76% loss of Hcrt-ir neurons (paired t test, t = 7.7; df = 2; p < 0.001; power = 94.6). ADA-ir neurons also showed a similar time course of neuronal marker loss. Representative photomicrographs from animals with unilateral Hcrt2-SAP lesions are presented in Figure 4A, B. The time course of loss of markers is consistent with that of other targeted saporin conjugates (Waite et al., 1994; Mantyh et al., 1997).

To identify whether the toxin affected other neuronal markers within the injection area, neurons containing melanin-concentrating hormone (MCH-immunoreactive) (Fig. 4C, D) or α-MSH (Fig. 5) were counted. MCH- and α-MSH-immunoreactive neurons are located in close proximity to the Hcrt neurons but are separate from the Hcrt neurons and also distinct from each other (Elias et al., 1998). MCH neurons were found to be decreased after unilateral Hcrt2-SAP injections (Figs. 3 and 4C, D) with the same time course as the Hcrt and TMN neurons, but α-MSH neurons were spared (Figs. 3 and 5). This suggests that some neurons are more sensitive to the toxin, and this sensitivity may depend on the presence of the Hcrt receptor and/or the subtype of Hcrt receptor on the neuron.

Experiment 3: Effects of bilateral injection of Hcrt2-SAP on sleep and wakefulness

The location of the injection sites is schematically illustrated in Figure 6. Figure 7 is a camera lucida drawing detailing the loss of the Hcrt-ir cells in two representative rats.

Extent of loss of Hcrt neurons

Animals administered bilateral injections of Hcrt2-SAP in the hypothalamus were compared with rats receiving injections of saline into the hypothalamus (n = 8; average number of Hcrt-ir cells per rat was 1324.0 ± 57.6). Animals administered bilateral injections of Hcrt2-SAP were divided into two groups: rats with >60% loss of Hcrt neurons (represented as Hcrt-x; n = 9; average number of Hcrt-ir cells per rat was 155.6 ± 55.6) and rats with <30% Hcrt cell loss (n = 3; average number of Hcrt-ir cells per rat was 928.0 ± 62.7). A one-way ANOVA identified a significant between-group difference (F(2,16) = 116.06; p < 0.001) and Tukey’s post hoc comparison revealed that all three groups were significantly different from each other with respect to the number of Hcrt-ir neurons (p < 0.001).

Hcrt2-SAP administered bilaterally to the LH produced a loss of Hcrt receptor mRNA (Fig. 8, A vs D) and of Hcrt-ir neurons (Fig. 8, B vs E). We estimate that the loss of Hcrt receptor mRNA-containing neurons extended along a radius of 0.8–1.0 mm, and this was sufficient to knock out the Hcrt-ir cells with boundaries extending from −2.0 to −4.0 posterior to bregma. Consistent with the loss of Hcrt-ir somata, there was a loss of Hcrt-ir fibers and terminals at target sites, such as the locus ceruleus (Fig. 8, C vs F).

Analysis of sleep data

Based on the time course of Hcrt neuronal loss after toxin administration (experiment 2 and Fig. 3), the sleep data obtained on days 2, 6, and 14 after injection were analyzed. There were no significant differences in sleep–wakefulness on day 2 after injection between the saline- and Hcrt2-SAP-treated rats. This indicates that both groups had similar sleep levels after the surgery and before day 4, when a noticeable cell loss was first evident in rats administered Hcrt2-SAP (Fig. 3). In the saline-treated rats, there were no significant differences in sleep between day 6 and day 14. Therefore, the data from these days were combined.

Figure 9 summarizes the diurnal distribution of sleep–wake states across the 24 hr in the three groups. As is typical of nocturnal rodents, saline-treated rats (n = 8) demonstrated a clear diurnal distribution of sleep–wake states, with the rats exhibiting more wakefulness at night and more sleep during the day. However, the Hcrt-x rats (n = 9) (Fig. 9A–C) demonstrated...
very little difference in sleep–wakefulness during the day versus during the night. Table 1 summarizes the ratio of sleep–wake states during the day versus during the night. In the Hcrt-x rats, the ratio for wakefulness and SWS is close to 1, indicating a lack of a diurnal difference in these states (Table 1). For REM sleep, the ratio is <1, indicating more REM sleep at night in the Hcrt-x rats (Table 1). The diurnal difference in EEG δ power (0.3–4 Hz) during periods of SWS was also attenuated in the Hcrt-x rats (Table 1).

The day versus night difference in wakefulness was lost because Hcrt-x rats exhibited 2.3 times as much total sleep time as controls during the dark period (p < 0.001). These rats had significant increases in both SWS (2.4 times; p < 0.001) and REM sleep (2.2 times; p < 0.001) at night. To determine how the overall percentage changes in sleep occurred, we determined the number and average duration of bouts of wakefulness, SWS and REM sleep during the day and night (Table 2). At night, the increase in sleep occurred because the duration of wake bouts decreased while the duration of SWS and REM sleep bouts increased (see Table 2 for significance). Regression analysis across all animals with lateral hypothalamic injections (n = 20) revealed a significant inverse relationship between the number of Hcrt-ir neurons and SWS (r = −0.84; df = 19; p < 0.01) and REM sleep (r = −0.74; df = 19; p < 0.01) during the dark period (Fig. 10).

During the day the Hcrt-x- and saline-treated rats had similar amounts of SWS. However, in the Hcrt-x rats sleep was significantly fragmented, as evidenced by a twofold increase in the number of transitions from SWS to waking (p < 0.01; Table 2). The increased awakenings during the day would prevent REM sleep from occurring (Mistlberger et al., 1987) and indeed, Hcrt-x rats had two-thirds fewer REM sleep bouts than saline-injected controls (p < 0.001; Table 2). Although these rats had significantly fewer REM sleep bouts, the lengths of individual REM sleep bouts were not different compared with saline rats (Table 2). Thus, during the day (the animal’s major sleep period), the sleep of Hcrt-x rats was highly fragmented, with frequent awakenings and fewer REM sleep bouts. At night, the Hcrt-x rats were awake less and slept more than saline-treated rats. When averaged over the 24 hr period, the Hcrt-x rats had significantly more SWS (36%...
increase) compared with saline-treated rats ($p < 0.01$) but had a slight (17%) decrease in the REM sleep amounts ($p < 0.05$). The emergence of the behavioral effects in the Hcrt-x rats corresponded with the time course of the loss of Hcrt-ir cells in the LH.

The day and night levels of sleep in rats with partial (30%) loss of Hcrt neurons were not different compared with saline-treated rats (Fig. 9D–F). However, when averaged over the 24 hr period, these rats had a small increase in SWS (6% increase; $p < 0.05$) and a decrease in REM sleep (26% decrease; $p < 0.01$) on day 14.

**SOREMPs**

SOREMPs are an important symptom of narcolepsy. Hcrt-x rats had on average $7.4 \pm 1.5$ SOREMPs per night and $3.2 \pm 1.2$ episodes during the day. Such episodes were rarely observed in saline-treated rats (day, $0.5 \pm 0.2$; night, $0.1 \pm 0.1$) or in rats with a $<30\%$ loss of Hcrt cells (day, $0.3 \pm 0.3$; night, $0.3 \pm 0.3$). A representative example of a normal REM sleep bout in a saline-treated rat versus a SOREMP in a Hcrt-x rat is presented in Figure 11. Representative examples of SOREMPs are evident in the video clips. The video recordings complement the data shown in Figure 11 by showing the behavioral repertoire exhibited by the rats before and after the SOREMPs. Moreover, the videos show the SOREMPs occurring at inappropriate times, including when the animals are feeding. Occasionally, the rats were found to exhibit “rocking” behavior as a prelude to a SOREMP (video 3).

When the rats entered into a SOREMP episode, such episodes lasted $2.1 \pm 0.2$ min on average, which is similar to the duration of REM sleep bouts (Table 2). Thus, the SOREMPs were inappropriate triggering of REM sleep during wakefulness.

The behavior of two rats was monitored until 45 d after injection, and an increase in sleep during the night (2.8 times increase in total sleep vs saline rats) and many SOREMPs were still evident (rats 171 and 181 in the video clips), suggesting a long-term change in sleep architecture. There was a significant negative correlation between the number of Hcrt cells in the LH and the number of SOREMPs during the night ($r = -0.72$; $df = 19$; $p < 0.05$).
Effects of loss of adenosine deaminase-immunoreactive neurons in the TMN on sleep

Hcrt2-SAP lesioned ADA-ir neurons in the TMN in many animals. The TMN neurons possess the Hcrt receptor (Fig. 2AB) but do not contain Hcrt. However, there was no significant relationship (p < 0.171) between the loss of ADA-ir neurons in the TMN and SWS or REM sleep. There was also no correlation between the number of adenosine deaminase-containing cells in the TMN and the number of SOREMPs.

DISCUSSION

This study demonstrated in rats that lesion of the LH, which also eliminated the Hcrt-containing neurons, increased both SWS and REM sleep and produced SOREMPs. The effects were site-specific, because the application of the saporin conjugate to a slightly more caudal site in the posterior hypothalamus (Fig. 6, filled triangles) did not produce hypersomnolence. The caudal site (TMN) contains the Hcrt receptor, but application of the saporin conjugate to this site lesioned the receptor-bearing neurons (Fig. 2) but did not produce narcoleptic-like sleep behavior.

Hcrt2-SAP

Saporin is a protein isolated from the seeds of Saponaria officinalis (Stirpe et al., 1983). Extensive studies have shown that when saporin is coupled with antibodies or ligands that recognize cell-surface antigens or receptors, the conjugate binding is specific and initiates apoptosis in targeted cells (Waite et al., 1994; Bergamaschi et al., 1996; Mantyh et al., 1997).

In this study, saporin was conjugated to the peptide hypocretin-2 (orexin-B). Prepro-hypocretin is cleaved into two smaller peptides, hypocretin-1 (orexin-A) and hypocretin-2 (orexin-B). Hypocretin-2 is a linear peptide with a free N terminus that facilitates conjugation to saporin. Hypocretin-1 is more difficult to couple to saporin because both of its termini are blocked (Sakurai et al., 1998). Hcrt2-SAP bound to cells containing the Hcrt receptor but did not bind to cells that did not contain the Hcrt receptor (KNRK cells), indicating specificity of the Hcrt2-SAP. Cytotoxic effects in the brain were demonstrated by administering the Hcrt2-SAP to LH neurons known to contain the Hcrt receptor. Specific markers of some neuronal phenotypes were decreased within the Hcrt2-SAP injection area, whereas others (a-MSH) were spared (Fig. 3).

Diurnal rhythm changes in the Hcrt-x rats

In the Hcrt-x rats, the diurnal rhythm of wakefulness and SWS was severely attenuated (Table 1), primarily because of an increase in sleep during the night. As a result, Hcrt-x rats had increased SWS (>36%) over the 24 hr period. Even rats with <30% Hcrt neuronal loss had a slight increase in SWS (>6%) over the 24 hr period. SWS and REM sleep were found to correlate with a decline in Hcrt neurons. Although the Hcrt-SAP-treated rats exhibited hypersomnolence, the 24 hr sleep of human narcoleptics is not different compared with normal controls (Aldrich, 1991). However, studies have not been done to specifically correlate sleep with the severity of Hcrt neuronal loss in human narcoleptics.

The diurnal rhythm of EEG δ power, a measure of sleep homeostasis (Borbely, 1994) was attenuated in the Hcrt-x rats (Table 1). However, there were no changes in the overall levels of δ power compared with saline-treated rats. δ power was not measured in Hcrt null mice (Chemelli et al., 1999). The Hcrt-x rats had an increase in both SWS and REM sleep during the normal night-active period, and daytime sleep was highly fragmented with frequent arousals. Human narcoleptics also exhibit excessive sleepiness and SOREMPs during the day, and nighttime sleep is very fragmented (Aldrich, 1991). Thus, in the Hcrt-x rats the sleep architecture is similar to what occurs in narcolepsy.

Because narcoleptics are sleepy during the wake–active period, it is hypothesized that in narcolepsy the circadian mechanism for arousal is impaired (Broughton et al., 1998). The diurnal rhythm of sleep–wakefulness is regulated by the suprachiasmatic nucleus (SCN), the circadian pacemaker. Lesions of the SCN eliminate the day–night variation in sleep, but such lesions have never been shown to produce SOREMPs (Cointet et al., 1975; Mistlberger et al., 1987). Thus, it is unlikely that the night–time hypersomnolence or the SOREMPs in the Hcrt-x rats were attributable to lesions of the SCN. In narcolepsy, the circadian clock functions normally (Dantz et al., 1994).

A possible explanation for the attenuation of the day–night sleep rhythm is that the lesion reduced the circadian drive from

Figure 6. Camera lucida drawings of injection sites as well as Hcrt-ir neurons (small dots) in the LH. Large filled circles represent the sites for which application of Hcrt2-SAP produced >60% Hcrt cell loss (Hcrt-x rats). Filled triangles represent the sites for which Hcrt2-SAP injections produced <30% Hcrt cell loss. Asterisks represent the saline injection sites. The location of Hcrt neurons (small dots) and ADA-ir neurons (marked by ×) is shown on the left side of the drawings. Arc, Arcuate nucleus; f, fornix; mt, mammillothalamic tract; PeF, perifornical area; TM, tuberomammillary nucleus; TMC, TM central portion. The nomenclature is according to the rat atlas of Paxinos and Watson (1986).
posterior hypothalamic arousal systems more likely is mediated through the medial preoptic and/or anterior hypothalamic nuclei (Abrahamson and Moore, 2001). We suggest that the SCN might regulate wakefulness by providing a waking signal to the Hcrt neurons. Such a signal might activate the Hcrt neurons, which then release Hcrt at target neurons to maintain wakefulness. We have recently proposed a model in which the Hcrt neurons activate the monoaminergic and cholinergic systems (Kilduff and Peyron, 2000). Intracerebroventricular injection of Hcrt induces arousal (Hagan et al., 1999), and systemic application of hypocretin-1 to narcoleptic dogs reduces cataplexy and normalizes sleep (John et al., 2000).

**SOREMPs**

As in the murine model, the Hcrt-x rats had more SWS and REM sleep at night and multiple periods of behavioral arrest during purposeful behavior. In the Hcrt knock out mice, EEG recordings were not made during the periods of behavioral arrest, whereas in the present study, continuous EEG recording allowed identification of the periods of behavioral arrest as SOREMPs (see video clips and Fig. 11). Interestingly, Shoham and Teitelbaum (1982), who made electrolytic lesions that included primarily the LH, observed that the rats collapsed into sleep when engaged in a spontaneous behavior such as grooming (they referred to such behavior as “groom-arrest”). Based on the present study, such groom-arrest episodes may very well have been SOREMPs.

Specific incidences of cataplexy were not observed in either hypocretin/orexin null mutant mice or the Hcrt2-SAP-injected rats. However, in canine narcolepsy, specific incidences of cataplexy are triggered in response to food or play and are short, lasting on the average 23 sec (Wu et al., 1999). Stimuli that trigger cataplexy need to be identified in rodents. Alternatively, the brainstem effector neurons implicated in cataplexy (Wu et al., 1999) need to be directly targeted to produce clear cataplectic episodes.

**Implications of these findings for narcolepsy**

In previous studies lesions of the posterior hypothalamus did not consistently produce symptoms of narcolepsy (Ranson, 1939; Nauta, 1946; Swett and Hobson, 1968; McGinty 1969; Danguir and Nicolaidis, 1980; Shoham and Teitelbaum, 1982; Sallan et al., 1988; Denoyer et al., 1991; Jurkowlaniec et al., 1994). Given the emerging evidence that narcolepsy is associated with a dysfunction or loss of the Hcrt system, at either the receptor or the ligand level, it is very likely that in previous studies the appropriate neurons were not destroyed.

There are also other neurons in the LH, and one can begin to assess their role in narcolepsy. For instance, MCH-containing neurons overlap with the Hcrt neurons (Elias et al., 1998), and these neurons were lesioned in the present study by Hcrt2-SAP. However, it is unlikely that loss of the MCH neurons could have caused the sleep abnormalities in our rats or human narcoleptics, because these cells are present in human narcoleptics (Thannickal et al., 2000) and MCH knock out mice do not show behavior consistent with narcolepsy (Shimada et al., 1998).

Loss of the histamine-containing neurons is also not the cause of the hypersomnolence or increased REM sleep, because the number of histamine-containing neurons was counted and there was no relationship with narcoleptic symptoms. The TMN has been implicated in the regulation of sleep–wakefulness because antihistamines cause drowsiness and the TMN contains the only
known collection of histaminergic neurons in the brain (Senba et al., 1985). The enzyme adenosine deaminase colocalizes with histamine and can be used to identify the histamine TMN neurons (Senba et al., 1985). Histamine microinjections into TMN targets such as the preoptic area produce a dose-dependent increase in wakefulness (Lin et al., 1994). The inhibition of histamine synthesis in the preoptic area increases sleep and decreases wakefulness (Lin et al., 1994). Histamine H1 and H2 receptors are postulated to mediate the arousal (Lin et al., 1994). TMN neurons have the highest discharge rate during waking and are

Figure 8. Photomicrographs of hypothalamic sections depicting Hcrt receptor mRNA (A, D), Hcrt-ir neurons (B, E), and Hcrt-ir fibers in the locus ceruleus (C, F). In A–C, tissue from saline-treated rats is depicted. D–F depict tissue from a representative rat with Hcrt2-SAP administered to the LH. A depicts an autoradiogram image of Hcrt receptor mRNA labeling in the LH (coronal section). The region outlined by the box in A represents the area in which Hcrt-ir neurons are present (B). Images in B and F are presented in reverse contrast. Hcrt2-SAP applied to the LH eliminated Hcrt receptor mRNA labeling (D) and Hcrt-ir neurons (E). Elimination of the Hcrt-ir neurons produced a loss of Hcrt-ir fibers at target sites such as the locus ceruleus (LC). In control rats, the LC is heavily innervated by Hcrt-ir fibers (C), but this innervation is lost after Hcrt2-SAP lesions of the Hcrt-ir neurons in the LH (F). 3V, Third ventricle; 4V, fourth ventricle; Amyg, amygdala; f, fornix; MHb, medial habenula; mt, mammillothalamic tract; PeF, perifornical nucleus.

Figure 9. Mean (±SEM) percentage of wakefulness, SWS, and REM sleep during 24 hr in rats administered Hcrt2-SAP or saline in the posterior hypothalamus. The 24 hr are represented in 2 hr blocks. The dark bar represents the 12 hr light-off period. Animals with a >60% decline in the number of Hcrt cells (A–C) experienced significantly more SWS and REM sleep at night compared with saline-treated rats. During the day the lesioned rats had as much SWS as controls but REM sleep was decreased. The night-time increase in sleep served to lessen the diurnal variation in sleep. In animals that had partial loss of Hcrt neurons (D–F), there was no change in sleep.
virtually silent during sleep (Vanni-Mercier et al., 1984; Szumylniak et al., 1989; Sakai et al., 1990). The TMN is a major target of the Hcrt-containing neurons. The Hcrt receptor mRNA is localized to the TMN (Fig. 2A,B) and Hcrt2-SAP lesioned the TMN. However, destruction of these neurons does not produce hypersomnolence, SOREMP, or other narcoleptic symptoms.

The Hcrt-x rats in the present study share many features of sleep architecture present in human (Aldrich, 1991) and canine (Kaitin et al., 1986a,b) narcolepsy and in Hcrt gene knock out mice (Chemelli et al., 1999). In humans (Aldrich, 1991), dogs (Lin et al., 1999), and the knock out mice (Chemelli et al., 1999), narcoleptic symptoms are not evident until adulthood, yet in our study with adult rats, a site-directed lesion readily produced these symptoms within a few days. This observation indicates that an inherited gene defect need not be the only route by which narcolepsy can occur. Humans without any familial history of the illness can develop narcolepsy, leading to the suggestion that this is an autoimmune disorder influenced by environmental factors (Honda and Matsuki, 1998). Increased number of astrocytes are present in the LH of narcoleptic patients (Thannickal et al., 2000), and a close association with a human leukocyte antigen DQ allele, DQB1*0602, is considered a predisposing factor in human narcolepsy (Mignot et al., 1997).

In the only two available animal models of human narcolepsy (Chemelli et al., 1999; Lin et al., 1999), the dysfunction in the Hcrt system is inherited and is in the entire animal. These characteristics make it difficult to localize the subgroup of Hcrt-containing neurons and the associated circuits that may underlie the individual symptoms of narcolepsy. The Hcrt2-SAP conjugate provides a method of investigating the contribution of the Hcrt system to the regulation of behavior across various species. This could be useful in determining whether the network underlying sleep is conserved across species. Moreover, because the Hcrt neurons project to multiple sites, Hcrt-SAP can be used to identify the role of these target sites in behavior. The effects of the saporin conjugate on sleep are long lasting, and this model could be used to test pharmacological treatments for narcolepsy.

**REFERENCES**


Figure 10. Relationship between sleep states (SWS and REM sleep) during the dark period and the numbers of Hcrt-ir cells in the LH. There was a significant inverse relationship between the numbers of Hcrt cells and sleep states.

Figure 11. Alternation between wakefulness (W), SWS, and REM sleep in rats administered saline (A) or Hcrt2-SAP (B) into the LH. The figure represents a 20 min segment of a sleep–wake recording during the night (9:00 P.M.). A and B consist of a recording of the EEG, power of the EEG in the δ (0.3–4 Hz; pink) and θ (4–12 Hz; yellow) bands, and integrated activity of the nuchal muscles (EMG). The sleep–wake state determination, based on the relationship of the EEG, power, and EMG activity, is indicated at the bottom of each panel. A depicts a normal transition from SWS to REM sleep to wakefulness. B depicts a SOREMP exhibited by a Hcrt2-SAP-treated rat with a 90% loss of Hcrt-ir neurons. The SOREMP is identified by a loss of EMG tone (near zero), by increased θ activity, by a reduction in δ activity (pink band in B), and by an EEG amplitude that is similar to wakefulness. These criteria are used to identify REM sleep, including SOREMP, and they are not present during wakefulness or SWS. Note that the first brief bout of wakefulness in B cannot be construed as REM sleep, because there is no θ activity and the EMG tone is rising, denoting that the rat woke up, albeit briefly.


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