Inhibition of Serotonergic Neurons in the Nucleus Paragigantocellularis Lateralis Fragments Sleep and Decreases Rapid Eye Movement Sleep in the Piglet: Implications for Sudden Infant Death Syndrome

Robert A. Darnall,1 Michael B. Harris,2 W. Hugh Gill,3 Jill M. Hoffman,1 Justin W. Brown,1 and Mary M. Niblock1

1Department of Physiology, Dartmouth Medical School, Lebanon, New Hampshire 03756, 2Institute of Arctic Biology, University of Alaska Fairbanks, Fairbanks, Alaska 99775-7000, and 3Tulane School of Medicine, New Orleans, Louisiana 70115

Serotonergic receptor binding is altered in the medullary serotonergic nuclei, including the paragigantocellularis lateralis (PGCL), in many infants who die of sudden infant death syndrome (SIDS). The PGCL receives inputs from many sites in the caudal brainstem and projects to the spinal cord and to more rostral areas important for arousal and vigilance. We have shown previously that local unilateral nonspecific neuronal inhibition in this region with GABA_A agonists disrupts sleep architecture. We hypothesized that specifically inhibiting serotonergic activity in the PGCL would result in less sleep and heightened vigilance. We analyzed sleep before and after unilaterally dialyzing the 5-HT_1A agonist (+)-8-hydroxy-2-(dipropylamino)-tetralin (8-OH-DPAT) into the juxtafacial PGCL in conscious newborn piglets. 8-OH-DPAT dialysis resulted in fragmented sleep with an increase in the number and a decrease in the duration of bouts of nonrapid eye movement (NREM) sleep and a marked decrease in amount of rapid eye movement (REM) sleep. After 8-OH-DPAT dialysis, there were decreases in body movements, including shivering, during NREM sleep; body temperature and heart rate also decreased. The effects of 8-OH-DPAT were blocked by local pretreatment with N-[2-(2-methoxyphenyl)-1-piperazinyl][ethyl]-N-2-pyridinylcyclohexane-carboxamide, a selective 5-HT_1A antagonist. Destruction of serotonergic neurons with 5,7-DHT resulted in fragmented sleep and eliminated the effects of subsequent 8-OH-DPAT dialysis on REM but not the effects on body temperature or heart rate. We conclude that neurons expressing 5-HT_1A autoreceptors in the juxtafacial PGCL are involved in regulating or modulating sleep. Abnormalities in the function of these neurons may alter sleep homeostasis and contribute to the etiology of SIDS.

Key words: serotonin; sleep; sudden infant death syndrome; 5-HT_1A receptor; REM; 8-OH-DPAT

Introduction

Sudden infant death syndrome (SIDS) remains the most common cause of death in postneonatal infancy. A major advance in our understanding of the etiology of SIDS has been the finding that many SIDS infants have decreased binding of muscarinic and kainate receptors in the arcuate nucleus at the ventral surface of the medulla (Kinney et al., 1995; Panigrahy et al., 1997) and abnormal binding to serotonergic (5-HT) receptors in the medullary serotonergic nuclei, including the paragigantocellularis lateralis (PGCL) (Panigrahy et al., 2000; Kinney et al., 2003). Dysfunction in these regions may increase the risk for SIDS by altering protective reflexes to stressors encountered during sleep such as hypercapnia, hypoxia, and laryngeal stimulation (Filiano and Kinney, 1994). In conscious animals, the local dialysis of GABA_A agonists into the PGCL decreases the ventilatory response to hypercapnia (Curran et al., 2001) and prolongs the laryngeal chemoreflex (Van der Velde et al., 2003).

Sleep represents a period of relative vulnerability, when many control systems function at lower levels. Although SIDS presumably occurs during sleep, there is little evidence to support a specific sleep abnormality. Sleep research in SIDS has been focused on arousal as an important protective mechanism (Kahn et al., 2002). However, an increase in arousability may also be disadvantageous and lead to fragmented sleep or sleep deprivation (Simpson, 2001). We have reported previously that nonspecific neuronal inhibition with GABA_A agonists in the PGCL region in the piglet disrupts sleep architecture and results in more wakefulness (WAKE) (Darnall et al., 2001).

The results of our previous experiments and the findings of altered serotonergic receptor binding in the PGCL in SIDS infants prompted us to look more closely at the role of PGCL 5-HT neurons in sleep. The PGCL extends caudally from the superior olive to the anterior pole of the lateral reticular nucleus (Andrezik et al., 1981). It receives inputs from the nucleus of the solitary tract, A1 region, parabrachial nucleus, Kolliker-Fuse nucleus, periaqueductal gray, and the hypothalamus. In addition, the...
more rostral (juxtafacial) PGCL receives polymodal sensory inputs from the inferior colliculus, the dorsal column nuclei, and the medial geniculate nucleus (Van Bockstaele et al., 1989, 1993; Van Bockstaele and Aston-Jones, 1995). The PGCL sends important projections to more rostral areas important for alertness and arousal, including the locus ceruleus (LC) (Aston-Jones et al., 1986, 1991a), and to both the dorsal and ventral horns of the spinal cord (Holstege and Kuyper, 1987) with extensive collaterali zation supplying multiple spinal cord segments (Bowker and Abbott, 1990; Kaus, 1991). Thus, neurons located in the PGCL are positioned to play an important role in integrating multiple sensory inputs for modulating brain and spinal alerting systems.

In this study, we focused on 5-HT neurons in the juxtafacial PGCL and asked whether decreasing serotonergic activity would alter normal sleep patterns. Our paradigm was to evaluate sleep after unilaterally decreasing serotonergic activity in the PGCL using the selective 5-HT1A agonist (+)-8-hydroxy-2-(dipropylamino)-tetralin (8-OH-DPAT).

Materials and Methods

Experiments were performed on piglets of either sex, aged 4 – 18 d and weighing 1.5 – 3.8 kg at the time of study. All surgery and experimental protocols were approved by The Institutional Animal Care and Use Committee of Dartmouth College. When not in the laboratory, the piglets were housed with the sow and siblings in a farrowing crate located in the animal care facility. Other investigators in the caudal brainstem (Berner et al., 1999), and previous observations showing that young newborn piglets sleep in bouts throughout the 24 h period with little consolidation during the dark cycle, all studies were performed between 10:00 A.M. and 3:00 P.M. and were of similar recording duration. The plethysmograph was sealed −1.5 h before the experiment to allow the temperature and humidity to stabilize. Calibration was performed using sequential triplicate injections of 1, 2, 3, and 5 ml of air. The piglet was then placed in the plethysmograph and connected to the monitoring equipment. The microdialysis probe was inserted and dialysis started with artificial CSF (aCSF) [containing the following (in mM): 152.2 Na, 3.0 K, 131.1 Cl, and 1.5 Ca, adjusted to a pH of 7.4] at a flow rate of 8.5 μl/min. After temperature, humidity, [CO2], and [O2] reached stable values (≈1 h), measurements were begun. Two protocols were used in this study: (1) an experimental protocol in which normal sleep cycling was first recorded for ≈2 h, during which aCSF was continuously dialyzed. The dialysate was then switched to either 10 or 30 mM 8-OH-DPAT (Sigma, St. Louis, MO) and continued for 30 min. The dialysate was then switched back to aCSF for the remainder of the experiment; and (2) a time-control protocol in which aCSF was substituted for the period of 8-OH-DPAT dialysis resulting in continuous aCSF dialysis throughout the experiment.

To confirm that our results were secondary to activating 5-HT1A receptors, four animals were dialyzed for 30 min with 30 mM 8-OH-DPAT after pretreatment for 30 min of local dialysis with N-[2-[4-(2-methoxyphenyl)-1-piperazinyl]ethyl]-N-2-pyridinylcyclohexane-carboxamide (WAY100635; Sigma), a selective, “silent” 5-HT1A-receptor antagonist. The term silent 5-HT1A-receptor antagonist has been used to distinguish true antagonists from partial agonists and describes compounds that lack intrinsic activity yet effectively block the effects of receptor agonists. To further determine whether our results were attributable to activating somato-dendritic 5-HT1A autoreceptors or postsynaptic 5-HT1A receptors, 5,7-dihydroxytryptamine (5,7-DHT; Sigma), a toxin selective for 5-HT neurons, was dialyzed into the PGCL of four animals after pretreatment with desipramine hydrochloride to prevent destruction of catecholaminergic neurons. After a week, 30 mM 8-OH-DPAT was dialyzed into the same region of the PGCL, and the results were compared with a group of animals of similar ages not treated with 5,7-DHT.

8-OH-DPAT doses. Relatively large doses of 8-OH-DPAT were used in the current study compared with those used in other dialysis experiments in the pons (Darnall et al., 1996). Keeping in mind that the estimated tissue concentration is 1/10 of the dialysate concentration (De Lange et al., 1995), 10 mM 8-OH-DPAT (estimated 1 mM tissue concentration) produced some effects on sleep, but 30 mM was necessary to obtain consistent results. These doses are consistent with those used by other investigators in the caudal brainstem (Berner et al., 1999), and larger doses may be necessary for a number of reasons. Compared with dorsal raphe 5-HT neurons, caudal medullary 5-HT neurons have faster firing rates (Heym et al., 1982b), may have fewer 5-HT1A autoreceptors (Trulson and Frederickson, 1987), and appear to be less sensitive to 5-HT1A agonists (Heym et al., 1982a). Our fluorescent data and 5,7-DHT data, although not conclusive, provide some level of confidence that we were affecting 5-HT1A receptors in an area restricted to the juxtafacial PGCL and a portion of the retrotrapezoid nucleus.

Data reduction and calculations. Data reduction, including sleep scoring, was done using custom programs written in Matlab (MathWorks, Natick, MA). For sleep-state scoring, a wavelet-based analysis that has been described previously (Darnall et al., 2001) was used to derive frequency information from the EEG. Slow wave (θ) activity was estimated by combining levels 6 – 9 of a 9 level discrete wavelet decomposition/reconstruction (0.3 – 4.7 Hz), and theta activity was estimated by using level 5 (4.7 – 9.4 Hz). A similar analysis was done for EOG and nuchal EMG recordings to isolate the most important features of each signal. Periods of non-rapid eye movement (NREM) sleep, rapid eye movement...
aged in bins and smoothed using a 60 s time constant. VO2 was for counting, on either side of the midline. The rectangle extended later-

al counters within an identical counting area on the unlesioned side. Rectanglesioned side within a standardized counting area and compared with

PGCL, TPOH-immunoreactive (TPOH-ir) cells were counted on the

exception that monoclonal mouse anti-rat tyrosine hydroxylase (TyrOH) were also identified using an identical protocol, with the

nohistochemistry in the piglet brainstem has been described in detail

the results section.

For cardiorespiratory variables, the original digitized data were resa-

mplied at rates appropriate for the variable. For respiratory calculations, the maximum and minimum of each breath related pressure fluctuation were determined using an automated peak detector followed by manual correction, if necessary. The amplitude of each breath (maximum − minimum) was used to derive breath to breath tidal volume (Vt) (Bar-

tlett and Tenney, 1970). Minute ventilation (V\text{\dot{E}}) was calculated as the product of Vt and instantaneous respiratory rate (RR) calculated from the interbreath interval. The peak of each blood pressure pulse was de-

termined similarly and was used to calculate beat-to-beat HR. BPm was calculated from the arterial pressure waveform.

For metabolic variables, body and plethysmograph temperature, inlet and outlet [O2], and plethysmograph gas flow rate were aver-

aged in 1 s bins and smoothed using a 60 s time constant. CO2 was calculated from the difference in the fractional inlet and outlet [O2] and the gas flow rate. VCO2 was similarly calculated assuming an inlet room air fractional concentration of 0.0003. We made no attempt to correct the measurements for water content or the respiratory quotient (RQ) and assumed no major changes in either over the course of the experiments.

Neuroanatomy. At the conclusion of experiments, each piglet was killed with an injection of sodium pentobarbital followed by an intracar-
diack injection of 5–10 ml of saturated potassium chloride. Microinjec-
tions of 20–50 μl of 1% potassium permanganate were made through a

bed microdialysis probe to mark the location of the tip of the microdialysis probe in reference to external landmarks (Sun et al., 2000). The brainstem was removed and frozen in cryoembedding medium (Tissue-Tek OCT; Sakura Finetek, Torrance, CA). Brainstems were cryo-

sectioned (40 μm) at −18°C, and sections were thaw-mounted on gelatin-coated glass slides. Sections were fixed for 10 min in 37%

phosphate-buffered formalin, pH 7.4, and then stained with cresyl violet. The rostrocaudal length of the brainstem differed among piglets over the course of the experiments, and coordinates expressed relative to the bregma interaxial line did not always accurately describe the location of dialysis probes with respect to internal medullary landmarks. Therefore, we expressed the location of each probe with respect to three relevant internal medullary structures: the midline, the ventral surface, and the caudal pole of the facial nucleus (Curran et al., 2001). For convenience, the lesions are plotted both in reference to the obex and the facial nucleus in the results section.

To determine the extent of the neuronal destruction caused by local

dialysis of 5,7-DHT, immunohistochemical methods were used to iden-
tify neurons containing tryptophan hydroxylase (TPOH). TPOH immu-
nohistochemistry in the piglet brainstem has been described in detail

previously (Niblock et al., 2004). Adjacent sections were stained with
cresyl violet for anatomical comparisons and identification of medullary nuclei and landmarks. In addition, neurons containing tyrosine hydroxylase (TyrOH) were also identified using an identical protocol, with the exception that monoclonal mouse anti-rat tyrosine hydroxylase anti-

body (Sigma) was used as the primary antibody.

To determine the extent of the destruction of 5-HT neurons in the

PGCL, TPOH-immunoreactive (TPOH-ir) cells were counted on the

lesioned side within a standardized counting area and compared with

counts within an identical counting area on the unlesioned side. Rectan-
gles of equal dimensions were superimposed onto each slice that was used for counting, on either side of the midline. The rectangle extended later-

dSpin 2-D analysis.

Results

Distribution of 5-HT neurons in the piglet medulla and

anatomic locations of dialysis probe tips

The distribution of TPOH-ir neurons in the piglet brainstem and its relationship to other species, including the human, has been described in detail previously (Niblock et al., 2004). In the piglet, the PGCL contains 5-HT neurons that lie in a column lateral to the midline that extends from the ponto-medullary border to several millimeters caudal to the caudal pole of the facial nucleus. The more rostral (juxtafacial) portion of the PGCL lies mostly medial to the facial nucleus. In the dorsoventral dimension, it

ally from 1 to 4.2 mm lateral to the midline and from the ventral surface to 3.1 mm dorsal to the surface at the most medial dimension. Every sixth

40 μm section was counted, extending from 18 sections caudal to the end of the lesion to 18 sections rostral to the most rostral edge of the lesion. In a few instances, there were areas of tissue destruction within the counting area caused by the dialysis guide tube. To correct for this, the mirror image of the damaged area was superimposed on the unlesioned side, and cells within this area were excluded from the cell count. Cells were counted only if they were within the counting area, morphologically identifiable as neurons, axon and dendrite(s) were visible, and the neu-

ronal cytoplasm had a dense distribution of reaction product that ex-

cluded the nucleus if visible.

To determine whether 5-HT1A receptors localize on 5-HT neurons in the

PGCL, neurons were double labeled with antibodies for TPOH and the

5-HT1A receptor. Forty micrometer sections were fixed in 4% para-

formaldehyde at 4°C for 30 min, blocked in 4% normal goat serum, and

incubated in primary antibody overnight at 4°C. Sections were washed in PBS with 0.1% Triton-X and incubated in appropriate anti-rabbit and anti-mouse fluorescent secondary antibodies (Alexa Fluor) for 2 h at room temperature. Sections were then washed in PBS with 0.1% Triton-X and allowed to dry for 60 min at room temperature before being coverslipped in Fluoromount-G (Southern Biotechnology, Bir-

mingham, AL).

Analysis and statistics. Twenty-one animals of either sex were studied

don different days to determine the effects of 8-OH-DPAT on sleep. Time

control experiments were performed on six animals, in which aCSF was
dialized for the entire study period. Six experiments were performed with 10 mx 8-OH-DPAT, and 17 experiments were performed using 30

mx 8-OH-DPAT. An additional four animals were studied with both

WAY106635 and 8-OH-DPAT, and the results were compared with both the

six control and 17 30 mx 8-OH-DPAT experiments. Another four animals were studied with 30 mx 8-OH-DPAT 1 week after dialysis of

5,7-DHT. The results of these experiments were compared with a sub-
group of piglets from the 30 mx 8-OH-DPAT experiments that were

≥12 d of age at the time they were studied.

Probe tips were considered to be in the appropriate position if there was a high likelihood that dialyzed 8-OH-DPAT would extend into the

PGCL. We therefore accepted probe tip locations that were at least 1 mm lateral to the midline and ≥1.5 mm lateral to the medial edge of the facia

nucleus. For the analysis, we treated each study day as a separate case.

Bout number, bout duration, and the percentage of time spent in each

state was determined for NREM, NREM, and WAKE, as well as slow wave or delta activity, integrated EEG amplitude, and integrated neck EMG activity. Data were averaged for the hour before and the hour starting

with the onset of 8-OH-DPAT (or sham aCSF) dialysis and entered into a

three-way ANOVA (Systat version 10.2; Systat, Evanston, IL). The dia-

lysat (aCSF or 8-OH-DPAT) and the state (NREM, REM, WAKE) were

considered repeated or within-subjects factors, and the 8-OH-DPAT
dose (0, 10, or 30 mx) was used as a grouping factor. In some analyses, age or a combination of age and weight was added as covariates in the

analysis. Post hoc tests were performed if there were significant interac-
tions between factors. Pairwise comparisons were done by computing a

critical T value from the MS error term (Winer, 1962) and correcting the

resulting probability for multiple comparisons. Values were expressed as

means ± SEM, and the criterion for statistical significance was set at p < 0.05.
8-OH-DPAT as determined by dialysis of 30 mM fluorescein (shown in yellow). The rationale for and problems associated with using fluorescein to determine distribution volume has been described previously (Curran et al., 2001). Nevertheless, the distribution of fluorescein provides a rough estimate of the extent of spread of dialedyzed compounds of similar molecular weights. A 30 min dialysis of 30 mM fluorescein produced a 3-D ellipse that was ~2.1 mm wide and ~3.5 mm long with a volume of 13.2 μl. This value is approximately twice as large as the volumes obtained in our laboratory using 10 mM fluorescein dialyzed for 20 min in chronically instrumented piglets (5.8 μl) (Messier et al., 2002) or 10 min in decerebrate piglets (6.3 μl) (Curran et al., 2000). A typical section at the level of the facial nucleus is shown in Figure 1B, illustrating the location of TPOH-ir neurons.

Figure 2A shows the location of the dialysis probe tips in the study animals referenced to the obex, and Figure 2B shows the rostrocaudal and mediolateral position of the probe tips in relation to a standardized facial nucleus as discussed in Materials and Methods. Note that most of the tip locations are clustered medial to the facial nucleus.

**Effect of 8-OH-DPAT on sleep architecture**

Our initial analysis using 0 mM (time control; aCSF), 10 mM, and 30 mM 8-OH-DPAT revealed significant interactions between dose, state, age, and the effect of 8-OH-DPAT on several measures of sleep architecture, including bout number, bout duration, and time spent in each state. However, the effect of 10 mM 8-OH-DPAT was not consistently different from that of aCSF alone. In contrast, the effects of 30 mM 8-OH-DPAT on all variables were consistently different from both the effects of aCSF and 10 mM 8-OH-DPAT. We therefore elected to report in detail only the effects produced with 30 mM 8-OH-DPAT dialysis. Six animals were in the control group and received continuous dialysis with aCSF. They weighed 2.50 ± 0.27 kg and were 8.3 ± 1.5 d of age. In 17 experiments, animals were dialyzed with aCSF for ~2 h and then were dialyzed with 30 mM 8-OH-DPAT for 30 min followed by aCSF dialysis for the remainder of the experiment. The animals in this group weighed 2.61 ± 0.12 kg and were 10.3 ± 0.8 d of age. There were no significant differences in the weights and ages of the two groups.

Because our studies were performed over an age range of 4–18 d, we first examined the relationship between age and measures of sleep architecture. Over the age range of the piglets that we studied, there was no relationship between age and bout number, bout duration, or the percentage of time spent in any state. However, the effect of 8-OH-DPAT on bout number, bout duration, and time spent in each state. However, the effect of 8-OH-DPAT on several measures of sleep architecture, including bout number, bout duration, and time spent in each state.

---

**Figure 1.** 3-D reconstruction of the piglet brainstem showing the distribution of 5-HT neurons. **A**, A cross-sectional view is on the left, and a longitudinal view is on the right. TPOH-ir neurons are shown as red dots. The blue shaded areas are the approximate locations of the facial nuclei. The yellow-green area is the approximate diffusion area of dialyzed 30 mM 8-OH-DPAT centered in the juxtafacial PGCL as determined by fluorescein. **B**, The left panel shows a section through the medulla near the caudal end of the facial nucleus (indicated by the dashed line in **A**). TPOH-ir neurons are stained brown. The rectangle contains 5-HT neurons located in the PGCL, which is in section lie lateral to the pyramids and medial to the facial nucleus. The right panel is an enlarged view of the area within the rectangle shown in the left panel.

---

**Figure 2.** Location of the dialysis probe tips. **A**, Location of the dialysis probe tips referenced to the obex. **B**, Location of the dialysis probe tips referenced to a standardized facial nucleus to take growth and different sizes of the animals into account. The black rectangle represents the standardized facial nucleus. Probe tips located within the rostrocaudal dimension of the nucleus are plotted as absolute distances from the midline and relative distances between the caudal and rostral pole of the facial nucleus. SOC, Superior olivary complex; NTS, nucleus of the solitary tract; IOG, inferior olive; pyr, pyramid; DHT, 5,7-DHT; VII, facial nucleus.
state. Age-related effects were taken into account in subsequent analyses when appropriate.

The main effect of dialyzing 8-OH-DPAT into the PGCL was fragmentation of sleep associated with a dramatic reduction in the amount of REM sleep. We defined sleep fragmentation as an increase in the number and a decrease in the duration of bouts of NREM sleep. Figure 3 shows data from a typical experiment. Note the regular cycling of sleep states before dialysis compared with the fragmented pattern during and shortly after 8-OH-DPAT dialysis. The most striking result was the marked decrease in the number of REM bouts and the percentage of time spent in REM after 8-OH-DPAT dialysis. REM was completely abolished in 53% (9 of 17) of the piglets. In the remaining eight piglets, there was a 75.4% decrease in the number of REM bouts. Figure 4 shows the effects of 8-OH-DPAT dialysis on bout number, bout duration, and the percentage of time spent in each state. Note the decrease in REM bouts associated with the increase in number of both NREM and WAKE bouts. In addition, bouts of NREM were shorter, resulting in no difference in the percentage of time spent in NREM. In contrast, bouts of WAKE were more frequent, but not of shorter duration, resulting in a significant increase in the percentage of time spent in WAKE.

In addition, the shorter periods of NREM were associated with lower levels of delta activity and integrated EEG amplitude after 8-OH-DPAT dialysis compared with continuous aCSF dialysis experiments. In addition, most animals exhibited a decrease in motor activity and a decrease in muscle tone during NREM after 8-OH-DPAT dialysis. Although an attempt was made to provide a thermoneutral environment, some animals exhibited shivering during NREM and WAKE during control periods, indicating that they were below their thermoneutral range. In these cases, there was a marked decrease or absence of shivering after 8-OH-DPAT dialysis. Neck EMG activity, on average, was also lower after 8-OH-DPAT dialysis. However, the small number of subjects accompanied by the large variation in the measurement of nEMG precluded a meaningful comparison between the changes in nEMG in the control (sham aCSF) group and the 8-OH-DPAT group. Mean and individual data are shown in Figure 5. Because of the possibility that periods that we scored as NREM after 8-OH-DPAT were examples of a dissociated state with REM-like hypotonia without rapid eye movements, we compared levels of neck EMG and EEG amplitude and δ activity during REM before 8-OH-DPAT with those during what we scored as NREM after 8-OH-DPAT. Although neck EMG amplitude, EEG amplitude, and δ activity were all lower during NREM after 8-OH-DPAT compared with control NREM values, they all remained considerably higher than during REM before 8-OH-DPAT (p < 0.03).

**8-OH-DPAT and the 5-HT₁₄ Receptor**

To confirm that the effects on sleep were secondary to activation of 5-HT₁₄ receptors, we locally dialyzed WAY100635, a selective

---

**Figure 3.** A typical 8-OH-DPAT dialysis experiment. Data include EEG, EOG, neck EMG, delta or slow wave activity (SWA), the TDratio, EEG amplitude, and a hypnogram showing NREM (filled black bars), REM (white bars), and WAKE (gray bars) periods. Periods of NREM sleep are characterized by increases in EEG amplitude and SWA activity, no rapid eye movements, and relatively increased nEMG. REM is indicated by low EEG amplitude and SWA, an increased TDratio, rapid eye movements, and low neck EMG amplitude. Transitions to WAKE are indicated by an abrupt increase in neck EMG, a decrease in the TDratio, and a gradual rise in SWA and EEG amplitude. **A,** Control experiment in which aCSF was dialyzed over the entire course of the experiment. The first panel is a period taken from the control period, and the second panel is taken from the sham experimental period, in this case continued aCSF dialysis. **B,** 8-OH-DPAT experiment. Recordings were taken from the baseline period (first panel) and experimental period (second panel), during which 8-OH-DPAT was dialyzed into the PGCL.

**Figure 4.** The effects of dialysis of 8-OH-DPAT (DPAT) into the PGCL on bout number per hour, bout duration, and the percentage of time spent in REM, NREM, and WAKE. In all panels, open circles are data from the control group (n = 6) in which aCSF was dialyzed during the entire experiment. Filled triangles are data from the 8-OH-DPAT group (n = 17) in which a period of aCSF dialysis was followed by a 30 min period of 8-OH-DPAT dialysis. The label “A” on the x-axis represents the mean of values recorded during a 1 h baseline period during which both groups were dialyzed with aCSF. The label “B” on the x-axis represents a 1 h experimental period during which the control group continued with aCSF dialysis and the 8-OH-DPAT group was dialyzed with 8-OH-DPAT. The differences between A and B in the control and 8-OH-DPAT groups were compared. An asterisk indicates a significant difference between the changes from A to B in the two groups. Error bars represent SEM.
5-HT<sub>1A</sub> receptor antagonist, into four animals before dialyzing 8-OH-DPAT. Figure 6 shows data from a single experiment. Note that sleep cycling is preserved during 8-OH-DPAT dialysis. The main effect of pretreatment with WAY100635, after taking age into account, was an inhibition of the effect of 8-OH-DPAT on REM bout number and percentage of time spent in REM. The ability of WAY100635 to prevent 8-OH-DPAT-induced changes in bout duration was less consistent. Figure 7 shows the group and individual data for bout number, comparing the WAY100635 data with both the control and 30 mM 8-OH-DPAT experiments. These data support the hypothesis that the effects of 8-OH-DPAT on sleep architecture are secondary to activation of 5-HT<sub>1A</sub> receptors.

We also hypothesized that the effects that we observed after 8-OH-DPAT dialysis were secondary to activation of 5-HT<sub>1A</sub> receptors located on 5-HT neurons and functioning as autoreceptors. Although it is widely accepted that 5-HT neurons located in the raphe colocalize with 5-HT<sub>1A</sub> receptors, there are no reports showing the presence of 5-HT<sub>1A</sub> receptors located on the somata or dendrites of 5-HT neurons in the PGCL. Although we did not do a quantitative analysis, we did confirm colocalization of TPOH-ir neurons and 5-HT<sub>1A</sub> receptors using immunohistochemistry. It was estimated that 90% of TPOH-ir neurons on each section also contained label for 5-HT<sub>1A</sub> receptors (D. S. Paterson, personal communication). The results from a representative section are shown in Figure 8. Examination of the distribution of the 5-HT<sub>1A</sub> receptor immunoreactivity suggests that 5-HT<sub>1A</sub> receptors are likely colocalized with both serotonergic and nonserotonergic neurons in the region of the PGCL that we studied.

To determine whether the effects that we observed were primarily secondary to activating autoreceptors, we unilaterally destroyed 5-HT neurons in the PGCL in four piglets by dialyzing 5,7-DHT, a selective toxin for 5-HT neurons, after pretreatment with desipramine to minimize destruction of any nearby TyrOH-containing neurons. After 1 week, 8-OH-DPAT was dialyzed into the regions, and the effects on sleep were evaluated and compared with the effects of 8-OH-DPAT dialysis in a subgroup of animals that were ≥12 d of age. Figure 9 illustrates the destruction of 5-HT neurons secondary to 5,7-DHT dialysis in one animal. Adjacent sections were stained with cresyl violet (Fig. 9A), for TPOH immunoreactivity (Fig. 9B), and for TyrOH immunoreactivity (Fig. 9C). Note the large area devoid of TPOH-ir neurons on the lesioned side (Fig. 9B). Also, there are very few TyrOH-immunoreactive (TyrOH-ir) neurons in this region, most lying more lateral to the rostral portion of the PGCL and importantly, as shown in Figure 9C, unaffected by 5,7-DHT. The cell counts are shown in Figure 10, showing the four individual piglets as well as the averaged data. The individual data show the location of the lesion in reference to the caudal pole of the facial nucleus, whereas the averaged data are referenced to the caudal edge of the lesion. The number of 5-HT neurons destroyed, expressed as a percentage of the control (nonlesioned) side, ranged from 39.7 to 77.5% and averaged 64.6 ± 8.7%.

By 1 week after 5,7-DHT dialysis, sleep was more fragmented.
The effects of 8-OH-DPAT on body temperature and HR CO₂ concentration in the plethysmograph did not change over outflow.

receptors located on nonserotonergic neurons that were involved compared with similar aged animals. There were more ($p = 0.023$) and shorter ($p < 0.05$) bouts of NREM per hour, with little or no change in the percentage of time spent in state. The bout number per hour, or bout duration of REM, was not different, and the percentage of time spent in REM was not affected. However, subsequent dialysis of 8-OH-DPAT did not decrease the amount of REM or change the relationship between the number, or duration, of bouts of WAKE or NREM, compared with a control group of similar ages (Fig. 11). Interestingly, the effect of 8-OH-DPAT on body temperature was not different in the animals treated with 5,7-DHT, compared with the control group of similar ages. Similarly, HR decreased after 8-OH-DPAT dialysis in the 5,7-DHT-treated animals similar to the changes in a similar-aged control group. These data indicate that the effects on sleep that we observed after 8-OH-DPAT dialysis were secondary to activation of somato-dendritic 5-HT₁A autoreceptors. However, the effects of 8-OH-DPAT on body temperature and HR may have been secondary to stimulating postsynaptic 5-HT₁A receptors located on nonserotonergic neurons that were involved in modulating these variables, perhaps by affecting sympathetic outflow.

Figure 7. Mean and individual bout number per hour data for REM, NREM, and WAKE from experiments in which WAY100635 was dialyzed before dialysis of 8-OH-DPAT (DPAT). Mean data from WAY100635 plus 8-OH-DPAT experiments (open squares; $n = 4$), control experiments in which aCSF was dialyzed over the entire study period (open circles; $n = 6$), and 8-OH-DPAT experiments in which aCSF dialysis was followed by 8-OH-DPAT dialysis (filled triangles; $n = 17$) are shown on the top panel. The label “A” on the x-axis represents the mean of values recorded during a 1 h baseline period during which all groups were dialyzed with aCSF. The label “B” on the x-axis represents a 1 h experimental period during which the control group continued with aCSF dialysis, the 8-OH-DPAT group was dialyzed with 8-OH-DPAT, and the WAY100635 plus 8-OH-DPAT group was first dialyzed with WAY100635 followed by 8-OH-DPAT dialysis. The differences between A and B in the control, 8-OH-DPAT, and WAY100635 plus 8-OH-DPAT groups were compared. An asterisk indicates a significant difference between the changes from A to B the 8-OH-DPAT group and the WAY100635 plus 8-OH-DPAT group. The control, 8-OH-DPAT, and WAY100635 plus 8-OH-DPAT groups are shown in the bottom panel plotted against age. Each point represents the difference between values during A and B for a single experiment. Error bars represent SEM.

The major findings are that dialyzing 8-OH-DPAT into the juxtafacial PGCL causes sleep fragmentation and a marked decrease in the amount of REM sleep. Although the rostral groups of 5-HT neurons play important roles in the regulation of sleep (Portas et al., 1996; Strecker et al., 1999; Monti et al., 2000; Sakai and Crochet, 2000; Sorensen et al., 2001), little is known about the role of 5-HT neurons in the PGCL in either regulating or modulating sleep. Large bilateral quisqualic acid lesions of the mediobulbar formation encompassing the juxtafacial PGCL decrease the amount of REM (Holmes and Jones, 1994). Nonspecific neuronal inhibition with GABAₐ agonists (Curran et al., 2001; Darnall et al., 2001; Messier et al., 2002) or lidocaine (Berner et al., 1999) in this region decreases the amount of sleep and increases wakefulness. In studies focusing on the role of 5-HT neurons in the midline raphe and immediate adjacent areas, dialysis or micro-injection of 8-OH-DPAT decreases sleep and increases wakefulness (Berner et al., 1999; Messier et al., 2004). We report the novel finding that 5-HT neurons in the juxtafacial PGCL play a role in the modulation of sleep and in particular, REM sleep. These findings suggest a possible link between sleep homeostasis and the abnormalities in serotoninergic receptor binding found in a large subset of SIDS infants in two large independent data sets.

The sleep effects of 8-OH-DPAT dialysis into the PGCL are caused by activation of 5-HT₁A somato-dendritic autoreceptors

It is likely that both somato-dendritic autoreceptors and postsynaptic 5-HT₁A receptors located on non-5-HT neurons were activated by 8-OH-DPAT in our studies (Thor et al., 1992; Kia et al.,...
However, the primary effects on sleep were caused by activation of 5-HT_{1A} somato-dendritic autoreceptors. This conclusion is based on our results showing that (1) there are large numbers of 5-HT neurons located in the PGCL in the piglet (Niblock et al., 2004), (2) most of these (~90%) colocalize with 5-HT_{1A} receptors, (3) pretreatment with the selective 5-HT_{1A} antagonist WAY100635 abolishes the sleep disrupting effects of subsequent 8-OH-DPAT dialysis, and (4) dialysis of 5,7-DHT into the PGCL produces sleep fragmentation similar to that produced with 8-OH-DPAT dialysis and eliminates the effect of subsequent 8-OH-DPAT dialysis on REM.

The effects of 8-OH-DPAT dialysis on body temperature and heart rate were likely attributable to the activation of postsynaptic 5-HT_{1A} receptors. 5-HT_{1A} receptors localize to nonserotonergic neurons in the raphe pallidus and parapyramidal region (Morrison, 2004) and in the rostral ventrolateral medulla (Holmes et al., 1994; Helke et al., 1997), where they are involved in thermoregulation and the control of blood pressure, respectively. Most of these neurons project to the intermediolateral column of the spinal cord and influence sympathetic activity involved in thermoregulation and cardiovascular control (Guyenet, 1990; Helke et al., 1997; Nakamura et al., 2004). Local application of 8-OH-DPAT into these regions, including the PGCL, in anesthetized animals consistently decreases blood pressure (Lovick, 1989; Helke et al., 1993). In contrast, we found in the conscious piglet that local dialysis of 8-OH-DPAT into the PGCL causes little or no change in blood pressure and a decrease in heart rate. These results are similar to those reported in two other studies in conscious animals in which 8-OH-DPAT was either dialyzed or microinjected into the caudal raphe and caused a decrease in heart rate with little or no change in blood pressure (Messier et al., 2004; Nalivaiko et al., 2005). Furthermore, destruction of 5-HT neurons with 5,7-DHT did not prevent the effects of subsequent 8-OH-DPAT dialysis on body temperature or heart rate, supporting the hypothesis that these effects were mediated by activation of postsynaptic 5-HT_{1A} receptors located on nonserotonergic neurons.

Activation of 5-HT_{1A} autoreceptors in the PGCL decreases the amount of REM sleep
A decrease in REM after 8-OH-DPAT dialysis in the PGCL was a consistent finding in all animals. Why does activation of 5-HT_{1A} autoreceptors in this region of the brainstem decrease REM sleep? One possibility is that sleep fragmentation after 8-OH-DPAT dialysis prevented the development of REM (Benington and Heller, 1994; Borbely, 1994). However, destruction of 5-HT neurons with 5,7-DHT did not prevent the effects of subsequent 8-OH-DPAT dialysis on body temperature or heart rate, supporting the hypothesis that these effects were mediated by activation of postsynaptic 5-HT_{1A} receptors located on nonserotonergic neurons.

Figure 8. Double labeling of TPOH-ir neurons and 5-HT_{1A}-immunoreactive receptors in one section of the PGCL near the caudal pole of the facial nucleus. A, TPOH-ir neurons are labeled in red. B, 5-HT_{1A} receptors are labeled in green. C, Overlay of A and B.

Figure 9. Unilateral destruction of TPOH-ir neurons on with 5,7-DHT. A, Standard Nissl stain (cresyl violet) showing many neurons scattered throughout the area. B, Section stained for TPOH-ir neurons. Note the absence of neurons on the lesioned (right) side. C, Section stained for TPOH-ir neurons. Note the scarcity of TyrOH-ir neurons in the PGCL on either side with their location limited to the very lateral edges of the section. Arrows point to the dialysis probe tip location. GC, Gigantocellularis; ROb, raphé obscurus; RPal, raphé pallidus; Pyr, pyramid; ION, inferior olive.

Figure 8.
surrounding regions might interrupt the normal state related attenuation leading to alterations in REM sleep homeostasis.

Sерotonergic neurons in the PGCL also have major projections to the dorsal horn of the spinal cord (Skagerberg and Bjorklund, 1985; Kwiat and Basbaum, 1992), where they modulate nociceptive (Basbaum and Fields, 1984; Leung and Mason, 1999) and non-nociceptive (Gray and Dostrovsky, 1983, 1985) sensory inputs. Furthermore, 5,6-DHT spinal cord lesions decrease REM in rats (Bjorkum et al., 1995), and intrathecal administration of 8-OH-5-DPAT increases the amount of sleep and decreases the amount of wakefulness (Bjorkum and Ursin, 1996). These data support the hypothesis that 5-HT neurons in the PGCL can modulate sleep state by dampening sensory input at the spinal cord level. Thus, inhibition of the activity of these neurons could result in an “undampening” of ascending sensory information promoting more wakefulness and the sleep fragmentation that we observed.

Dialysis of 8-OH-5-DPAT into the PGCL decreases muscle tone during NREM sleep
Skeletal muscle hypotonia during NREM similar to that seen during REM under control conditions was observed in some animals after 8-OH-5-DPAT dialysis. In a few animals, an attenuation in shivering during NREM was also observed, similar what occurs during REM under control conditions (Parmeggiani and Rabini, 1967). The classic studies by Magoun and Rhines suggested that neurons in the caudal medulla are involved in the modulation of

**Figure 10.** Cell counts showing the number of TPOH-ir neurons destroyed with 5,7-DHT on the lesioned side compared with the nonlesioned side. The top four panels show the counts from the individual animals referenced to the caudal pole of the facial nucleus (VII) to show the actual location of each lesion. Each set of points represent counts from one 40 μm section. Approximately every sixth section was counted. Open circles are counts from the unlesioned side, and filled circles are counts from the lesioned side. The bottom panel shows the group counts expressed as mean ± SEM referenced to the caudal end of the lesion.

**Figure 11.** Mean and individual data for bout number per hour for animals dialyzed with 5,7-DHT (DHT) and subsequently dialyzed with 8-OH-5-DPAT (DPAT) 1 week later compared with a group of animals of similar ages dialyzed with 8-OH-5-DPAT. The top panel shows the mean data. Open squares are the 5,7-DHT-treated animals (n = 4), and the filled triangles are a subgroup of 8-OH-5-DPAT experiments in which the animals were 12 d of age (n = 6). The label “A” on the x-axis represents the mean of values recorded during a 1 h baseline period during which both groups were dialyzed with aCSF. The label “B” on the x-axis represents a 1 h experimental period during which both the 5,7-DHT-treated and nontreated animals were dialyzed with 8-OH-5-DPAT. The differences between A and B in the 5,7-DHT-treated and nontreated groups were compared. An asterisk indicates that there is a significant difference between the changes in the two groups. In addition, # indicates that the baseline values were different in the two groups. Error bars represent SEM.

**Table 1. The effects of 8-OH-5-DPAT dialysis on selected metabolic variables**

<table>
<thead>
<tr>
<th>Variable</th>
<th>Control group</th>
<th>Experimental group</th>
<th>8-OH-DPAT</th>
</tr>
</thead>
<tbody>
<tr>
<td>VO₂ (ml·kg⁻¹·min⁻¹)</td>
<td>16.0 ± 0.8</td>
<td>13.3 ± 0.6</td>
<td>15.4 ± 0.5</td>
</tr>
<tr>
<td>VCO₂ (ml·kg⁻¹·min⁻¹)</td>
<td>13.3 ± 0.6</td>
<td>13.9 ± 1.0</td>
<td>13.2 ± 0.8</td>
</tr>
<tr>
<td>RQ</td>
<td>0.96 ± 0.05</td>
<td>1.15 ± 0.12</td>
<td>0.89 ± 0.04</td>
</tr>
<tr>
<td>Air temperature (°C)</td>
<td>26.6 ± 0.6</td>
<td>26.7 ± 0.4</td>
<td>26.3 ± 0.3</td>
</tr>
<tr>
<td>Body temperature (°C)</td>
<td>38.4 ± 0.2</td>
<td>38.5 ± 0.3</td>
<td>38.2 ± 0.2</td>
</tr>
</tbody>
</table>

Data include VO₂, VO₂, VO₂, and plethysmograph air temperature and piglet core body temperature. Dialysis of 8-OH-5-DPAT caused a small but significant (*) decrease in body temperature.

muscle tone (Magoun, 1944; Magoun and Rhines, 1946). Neurons in PGCL project to thoracic, lumbar, and sacral segments of the spinal cord (Kausz, 1991) and have axon collateral projections to both autonomic and somatic cell groups (Allen and Cechetto, 1994). Spinally projecting glycinerergic neurons that receive glutamatergic, cholinergic, and orexinergic inputs in this region have been implicated in the hypotonia associated with REM (Kodama et al., 1998; Lai et al., 1999; Hajnik et al., 2000; Mileykovskiy et al., 2002), and some appear to be serotonergic (Fort et al., 1993; Stornetta et al., 2004). It is possible that activation of either 5-HT₁₅ somato-dendritic receptors on glycinerergic/5-HT neurons or postsynaptic receptors located on non-5-HT glycinerergic neurons might result in a change in muscle tone.

What we scored as short periods NREM after 8-OH-5-DPAT dialysis might represent a dissociated state with REM-like hypotonia. However, there were no rapid eye movements or REM-like changes in respiration, blood pressure, and heart rate. In addition, although the levels of δ power, EEG amplitude, and neck EMG activity were lower during NREM after 8-OH-5-DPAT dialysis, they were significantly higher than during REM bouts before
8-OH-DPAT. We therefore believe that the relative hypotonia after 8-OH-DPAT dialysis into the PGCL is more likely related to the role of 5-HT neurons of providing tonic state related motor facilitation important for the modulation of muscle tone and rhythmic motor activity (Jacobs and Fornal, 1999).

Implications for sudden infant death syndrome

Victims of sudden infant death syndrome presumably die during sleep. However, there has been little evidence to support a specific sleep abnormality. The focus of research has been on protective arousal mechanisms that help defend against stressors often encountered during sleep. However, an increase in arousability may not always be an advantage because it might lead to fragmented sleep and sleep deprivation. Prolonged deep periods of recovery sleep might also increase the risk of obstructive sleep apnea (Simpson, 2001). REM sleep deprivation in adult humans and animals has been found to cause many behavioral changes (Dement, 1960; Koller et al., 1969), and brief periods of sleep deprivation in infants alters the autonomic control of heart rate (Franco et al., 2003) and decrease arousals (Franco et al., 2004). Our new findings suggest that abnormalities in medullary serotonergic neurons produce sleep fragmentation and decreased amounts of REM sleep. 5-HT$_{1A}$ receptor activation in this study was restricted to a region where abnormal serotonergic receptor binding has been demonstrated in SIDS infants. More widespread serotonergic receptor dysfunction may lead to considerable alterations in sleep homeostasis, increasing the risk for sudden death.

References


Helke CJ, Capuano S, Tran N, Zhuo H (1997) Immunocytochemical studies of the 5-HT1A receptor in ventral medullary neurons that project to the intermediolateral cell column and contain serotonin or tyrosine hydroxylase immunoreactivity. J Comp Neurol 379:261–270.


Kausz M (1991) Arrangement of neurons in the medullary reticular forma-
tion and raphe nuclei projecting to thoracic, lumbar and sacral segments of the spinal cord in the cat. Anat Embryol 183:151–163.


