Activity of Medial Mesopontine Units during Cataplexy and Sleep-Waking States in the Narcoleptic Dog

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Narcolepsy has been hypothesized to be a disease of rapid eye movement (REM) sleep. According to this hypothesis, cataplexy is a result of the triggering during waking of the mechanism that normally serves to suppress muscle tone in REM sleep. REM sleep control mechanisms have been localized to the pons. Narcoleptic dogs have increased numbers of cholinergic receptors in the medial pons. These findings suggest that neurons mediating the triggering of cataplexy might be located in medial pontine regions. In the present study, this hypothesis has been investigated by recording the discharge of units in the medial mesopontine region of the narcoleptic dog. Unit activity was examined in the nucleus reticularis pontis oralis, caudalis, and central gray, with each cell being recorded during both cataplexy and sleep states. Maximal discharge rates were observed, in all of these regions, during active waking states (mean rate, 45.3/sec) and REM sleep (16.0/sec), with minimal discharge rates in non-REM sleep (8.3/sec). Unit discharge was reduced in cataplexy relative to precataplexy periods. Cataplexy discharge rates were 8.3/sec, 52% of the mean REM sleep rate. Cataplexy discharge rates were also significantly lower than those at REM sleep onset. Cataplexy discharge rates were comparable to rates in quiet waking and non-REM sleep.

While medial mesopontine neurons discharge at high rates in REM sleep, they have little or no activity in cataplexy. We interpret the lack of activation of medial mesopontine units in cataplexy as indicating that the characteristic phasic motor activation of REM sleep does not occur in this state. We found that eye movements were significantly reduced in cataplexy relative to eye movements in waking periods preceding cataplectic attacks and relative to eye movements at REM sleep onset and during REM sleep.

We hypothesize that cataplexy is a result of the pathological waking activity of neurons that are normally selectively active in relation to the muscle tone suppression of REM sleep, combined with the sudden reduction of activity in brainstem neurons that are active in both REM sleep and waking. We have found REM-cataplexy-on cells in the medial mesopontine region also induces atonia (Lai and Siegel, 1988, Mitler et al., 1987). However, while cataplexy and sleep paralysis have a loss of muscle tone similar to that seen in REM sleep, these states are distinct from REM sleep in that consciousness is preserved (Guilleminault, 1976). It is unclear if other aspects of REM sleep physiology, such as altered autonomic activity, changes in thermoregulation, respiration, and phasic motor activation (Siegel, 1989) are present in cataplexy, or if atonia is the only common element.

The development of the narcoleptic dog model makes it possible to investigate these questions at the neuronal level. The narcoleptic dog presents dramatic episodes of cataplexy elicited by food presentation and play. Canine cataplexy is altered by pharmacological manipulations in a way that parallels the human condition (Baker and Dement, 1985; Miller et al., 1987; Mignot et al., 1989). As in human narcolepsy, cataplexy develops postnatally in dogs and shows some improvement with age. Also, as in the human condition, canine cataplexy is genetically determined (Honda et al., 1984; Baker and Dement, 1985; Mitter et al., 1987).

Studies of narcoleptic dogs and narcoleptic humans have found a marked upregulation of cholinergic (ACh) receptors in the medial mesopontine region (Boehme et al., 1981; Kilduff et al., 1986; Aldrich et al., 1990). Injection of cholinergic agonists in the medial mesopontine reticular formation regions of intact cats produces a loss of muscle tone without loss of consciousness that resembles cataplexy (Miller and Dement, 1974) or a REM sleep-like state (George et al., 1964; Van Dongen, 1978; Ka-tayama et al., 1984; Baghdoyan et al., 1987) depending on the exact injection site. Recent work in acute preparations has shown that microinjection of glutamate and corticotropin-releasing factor in the mesopontine region also induces atonia (Lai and Siegel, 1988, 1992). The receptor upregulation combined with the loss of muscle tone with chemical stimulation of this area suggests that it may be activated during both cataplexy and REM.
sleep. In the present study, we have recorded from cells in medial mesopontine and adjacent regions during sleep-waking states and cataplexy.

Materials and Methods

Three narcoleptic dogs, a female Labrador Retriever, a male Doberman Pinscher, and a female Doberman Pinscher, were bred at the Stanford University narcoleptic dog colony and transported to Los Angeles at the age of 8 months. Studies were performed over the next 3-10 month period. In an initial surgery, magnetic resonance (MR)-compatible high-nickel stainless-steel needles (Luftkin et al., 1987; Dackwiler et al., 1990) were implanted over the mesopontine reticular formation and mounted to the skull with Teflon screws and acrylic. The scalp was closed and MR imaging in the sagittal, coronal, and horizontal planes was performed under thiopental anesthesia. All scans were performed on a Fonar B-3000 M (Melville, NY) MR machine with a 0.3 tesla hybrid magnet utilizing a 256 x 256 matrix with 4.7 cm slice thickness and 1.28 pixels/mm. Spin echo pulse sequences were used to produce T1 weighted images with a repetition time of 500 msec and an echo time of 28 msec. The MR scans were then used to adjust stereotaxic coordinates using the implanted cannulas and the posterior limit of the inferior colliculus as reference points. The use of MR scanning was necessary because of the variability of skull size and resulting stereotaxic location of mesopontine targets. The correspondence between the stereotaxic atlas (Lim et al., 1960) and actual brain coordinates was found to be poor, with 1-5 mm deviations from the atlas in all three planes.

Units were localized to the nucleus reticularis pontis caudalis and oralis of the medial pons, corresponding to the FTG and FTL in the Berman (1968) nomenclature and to the central gray (Fig. 1). The discharge rates of medial mesopontine units located outside of the central gray did not differ significantly from those of central gray units (Table 1).

All units encountered had spontaneous activity in waking, with maximal discharge rates during movement, as we have previously seen in the analogous mesopontine regions of normal cats (Fig. 2) (Siegel and Tomaszewski, 1983; Siegel et al., 1983). Waking discharge was associated with directionally specific movements of the head and neck, as in the cat. Also, as in the cat, these units had high discharge rates in REM sleep and relatively low discharge rates in non-REM sleep (Figs. 3, 4). Cataplexies typically occurred during active waking periods and invariably were accompanied by an abrupt reduction in discharge rate. Figure 5 shows the activity of a pair of simultaneously recorded units during cataplexy. Unit activity and eye movements were reduced during cataplexy relative to pre- and postcataplexy periods. Cataplexy durations ranged from 12-74 sec (mean, 30.2 ± 14.9 sec). All units were recorded during cataplexies following phystostigmine administration and eight of the units were also recorded during cataplexies occurring without prior drug treatment. Unit discharge rates in cataplexies under the two conditions were not significantly different; the mean rate during cataplexies induced after phystostigmine treatment was 8.3/sec, while the rate during cataplexies induced without drug treatment was 11.1/sec (p > 0.5, paired t test). For purposes of statistical comparison, the 49 cataplexies induced after phystostigmine were used.

Discharge rate varied with arousal state (p < 0.001, two-way ANOVA). During elicited and spontaneous cataplexy, mesopontine units significantly decreased discharge relative to the immediately preceding waking rate (p < 0.05, Duncan's post hoc multiple range test). Cataplexy rates were significantly lower than REM sleep rates (p < 0.05, Duncan) and were comparable to non-REM and quiet waking rates. We saw no units selectively active in cataplexy or with increased activity in both cataplexy and REM sleep.

Figure 1. Plot of recorded cells at coronal levels RO to R1O from the Lim et al. (1960) atlas. RFC, Nucleus reticularis pontis caudalis; SO, superior olive; PT, pyramidal tract; IC, inferior colliculus; RPO, nucleus reticularis pontis oralis; SC, superior colliculus; PAG, periaqueductal gray; AQ, cerebral aqueduct; P, nucleus pontis.
PONTINE CAT-OFF CELL

Figure 2. Polygraph record of pontine cell showing reduction in discharge rate with cataplexy. Discharge rate is maximal in active waking and REM sleep, and intermediate in non-REM sleep. SM EEG, Sensorimotor cortex electroencephalogram; PL, posterolateral cortex electroencephalogram; EOG, electrooculogram; EMG, nuchal electromyogram; UNIT, pulse output of window discriminator triggered by unit.

Cataplexy intervals are considerably shorter than REM sleep periods. Since mean REM sleep discharge rates of mesopontine units are greater than their rates at REM sleep onset, we tested for the possibility that cataplexy rates might be similar to REM sleep onset rates. We found that medial mesopontine unit discharge rates were significantly lower in cataplexy than those in the same duration interval at REM sleep onset ($p < 0.01$, two-tailed paired $t$ test). Thus, medial mesopontine unit activity in cataplexy is not like REM sleep activity, when compared to REM sleep onset or the entire REM sleep period. We also found that the eye movement rate in cataplexy was significantly lower than that at REM sleep onset (Fig. 7; $p < 0.01$, two-tailed paired $t$ test).

We examined patterns of interspike intervals to determine if there might be some similarity in cataplexy and REM sleep pattern. However, we found that discharge patterns, as assessed by interspike interval histograms and autocorrelograms, showed no clear correspondence in cataplexy and REM sleep, even in those cells with discharge rates in cataplexy high enough to allow the calculation of meaningful spike train statistics. In general, patterns of interspike intervals tended to be less regular in cataplexy than REM sleep.

Table 1. Discharge rates of mesopontine cells in sleep-waking states and cataplexy ± SE ($n = 49$)

<table>
<thead>
<tr>
<th></th>
<th>QW</th>
<th>AW</th>
<th>Cataplexy</th>
<th>REM</th>
<th>non-REM</th>
</tr>
</thead>
<tbody>
<tr>
<td>All cells ($n = 49$)</td>
<td>8.8 ± 1.8</td>
<td>53.3 ± 8.2</td>
<td>8.3 ± 1.5</td>
<td>16.0 ± 3.5</td>
<td>8.3 ± 1.4</td>
</tr>
<tr>
<td>Central gray ($n = 18$)</td>
<td>7.5 ± 3.2</td>
<td>39.5 ± 12.6</td>
<td>5.8 ± 2.0</td>
<td>9.8 ± 3.1</td>
<td>4.5 ± 1.5</td>
</tr>
<tr>
<td>RPO-RPC ($n = 31$)</td>
<td>9.5 ± 2.1</td>
<td>48.6 ± 10.8</td>
<td>9.8 ± 2.1</td>
<td>19.6 ± 5.2</td>
<td>10.5 ± 1.9</td>
</tr>
</tbody>
</table>

QW, quiet waking; AW, active waking; RPO-RPC, nuclei reticularis pontis oralis- reticularis pontis caudalis.
Figure 3. Rate diagram indicating discharge rate as a function of state of cells plotted in Figure 1. OW, Quiet waking; AW, active waking; CAT, cataplexy; REM, REM sleep; NREM, non-REM sleep.

interspike interval histograms and autocorrelogram spike distributions were a function of rate, so that cataplexy patterns most closely resembled those of non-REM and quiet waking states.

**Discussion**

We found that medial mesopontine neurons in the narcoleptic dog have significantly reduced activity in cataplexy relative to both mean REM sleep and active waking rates. Discharge rates fell sharply at cataplexy onset from those present in the pre-cataplexy period. Cataplexy rates were also significantly lower than rates at REM sleep onset.

The majority of medial mesopontine neurons seen in this and previous studies discharge in relation to head, neck, thoracic, and related movements in waking (Rose, 1978; Siegel, 1979; Siegel and Tomaszewski, 1983; Siegel et al., 1983; Ni et al., 1990). We have interpreted their burst-pause discharge pattern in REM sleep as indicating that head, neck, and other axial movements are being commanded in this state, with their expression blocked by motoneuron hyperpolarization (Chase and Morales, 1989). This interpretation is supported by the release of such motor activities after lesions disrupting the system responsible for motor neuron hyperpolarization in REM sleep (Jouvet and Delorme, 1965; Henley and Morrison, 1974). It would follow from the above, and from our present findings, that central motor activation in cataplexy is reduced from precataplexy and REM sleep levels.

Our eye movement data support this interpretation. Eye movements decrease from prior waking levels at cataplexy onset. Eye movement levels in cataplexy are significantly lower than those in REM sleep and at REM sleep onset. These results, however, are somewhat surprising since one might expect attempted movements from the conscious paralyzed animals. Perhaps the expression of any such "willed" movements is blocked before it reaches mesopontine levels. Such a blockade or disfacilitation might have a role in the triggering of cataplexy. However, our previous work showing increased activity in a subpopulation of REM sleep active neurons in the medial medulla during cataplexy (Siegel et al., 1991) suggests that an active inhibitory mechanism is at least partially responsible for cataplexy.

Recent work in our laboratory in the decerebrate preparation has demonstrated that both atonia and motor activation are generated by colocalized pontomedullary systems, including the pedunculopontine/laterodorsal tegmental nuclei (PPN/LDT) and reticularis pontis oralis (RPO) regions, thus providing a mechanism for the generation of the "paradoxical" combination of muscle atonia with motor activity that characterizes REM sleep (Lai and Siegel, 1990). Further work has shown that non-NMDA glutamate receptors mediate the muscle tone suppression, while NMDA receptors mediate the motor activation elicited from atonia sites (Lai and Siegel, 1991). The presence of atonia without motor activation in cataplexy suggests a functional dissociation of the NMDA-mediated motor activation system from the non-NMDA-mediated atonia system.
Of particular relevance to the present work is our finding that stimulation of the colocalized atonia and locomotor control mechanisms leads first to atonia and then to locomotor activity with a time lag ranging between 10 and 40 sec (Lai and Siegel, 1991). This is the same sequence seen in REM sleep, where muscle tone suppression precedes the activation of mesopontine units and correlated rapid eye movements. We hypothesize that in short-duration cataplexies, the atonia mechanism is activated, as at REM sleep onset, but the recruitment of the NMDA-mediated phasic motor activation system is delayed. In extend-
ed cataplexy periods, a gradual transition to REM sleep may occur (Guilleminault, 1976), including not only rapid eye movements, but also the loss of consciousness of the outside world that characterizes sleep. Thus, the activation of the REM sleep phasic motor mechanism during the atonic state either is causally related to the loss of consciousness or, perhaps more likely, is correlated with the spread of excitation to other brainstem systems whose activity is linked to loss of consciousness.

Because of the absence of cataplexy-on cells in the medial pons, we hypothesize that a neuronal subpopulation in the PPN/LOT and lateral RPO atonia/locomotor control regions may be responsible for atonia in both REM sleep and cataplexy. According to this hypothesis, a cell population with activity selective for REM sleep and cataplexy should exist in these areas.

Our recent studies in the medial medulla support the idea that the atonia of cataplexy and REM sleep is mediated by a specialized subpopulation of cells. Most medial medullary cells, like the mesopontine cells recorded in the present study, were inactive in cataplexy but active in REM sleep (Siegel et al., 1991). However, we also found a group of medial medullary cells that were selectively active in cataplexy and REM sleep. These cells were localized to the ventral portions of nucleus magnocellularis, a region implicated in the suppression of tone in REM sleep (Sakai, 1980; Liu and Siegel, 1988; Schenkel and Siegel, 1989). These cells may be driven from a similar population of cataplexy-REM sleep-on pontine cells that have not yet been observed. One might predict the existence of such cells in the dorsolateral pons because REM sleep-on cells with projections to the medial medulla have been found in this region (Sakai, 1980; Shiromani et al., 1987). However, it is also possible that the medullary "cataplexy-on" cells do not require mesopontine excitation to produce the pathological loss of muscle tone in cataplexy, perhaps being activated autochthonously or with facilitation from adjacent medullary autonomic regions (Siegel et al., 1989). Thus, the determination of the activity pattern of pontine "REM sleep-on" cells in cataplexy is critical to an identification of the circuit producing this state.

Further examination of the differences and similarities between cataplexy and REM sleep will not only shed light on the mechanism responsible for narcolepsy, but will also provide important information on the mechanisms responsible for the suppression of muscle tone in REM sleep and the control of muscle tone in relation to locomotion in waking states.

References


