SUMMARY AND CONCLUSIONS

1. A total of 14.4% of medial reticular formation (RF) cells discharged maximally in relation to movements of the facial musculature. Cells related to movement of the ipsilateral eyelids, ipsilateral lip, ipsilateral pinna, and cells related to movement of several facial regions were seen.

2. Cells related to ipsilateral eyelid closure constituted 2.2% of all RF cells recorded. Responses to stroboscopic visual stimuli causing eyelid closure were seen at a 30- to 40 ms latency.

3. Cells related to ipsilateral lip movement constituted 2.2% of all RF cells recorded.

4. Cells related to ipsilateral pinna movement constituted 7.1% of all RF cells recorded.

5. Ipsilateral eyelid, lip, and pinna movement cells were completely silent during movements restricted to the contralateral facial musculature. They discharged only during movement of the affected ipsilateral region in a single direction. They were significantly more likely than adjacent nonfacial cells to respond during the reflex head shake and significantly less likely to respond to auditory stimuli. They had little or no tonic activity during waking and sleep.

6. Cells related to several facial movements constituted 2.9% of all RF cells recorded. Maximal discharge rates occurred during movement of the ipsilateral pinna, but smaller rate increases were also seen during movement of the ipsilateral lip and eyelid and during contralateral facial movements. In contrast to other facial cells, these cells also: a) responded during movements of ipsilateral facial regions in any direction, b) responded to somatosensory stimulation and passive movements of facial skin, c) responded to auditory and electric pulse stimulation of the skin, d) were inactive during the reflex head shake, and e) had high levels of tonic activity in both waking and sleep.

7. Each facial movement-related cell type had a different anatomical distribution within the brain stem. Cells related to eyelid, lip, and multiple facial movements formed localized clusters ventral to the abducens nucleus. Cells related to pinna movement were scattered throughout the gigantocellular tegmental field. Facial movement-related reticular cells may have premotor and proprioceptive roles in the regulation of facial movements.

INTRODUCTION

Facial movements in the cat are controlled by at least 35 muscles, 20 of which insert on portions of the auricle (3). The motoneurons controlling these muscles are contained in the nucleus of the 7th nerve. The facial nucleus is known to receive projections from the region around the 3rd nerve nucleus, the midbrain paralaminar region, the parabigeminal region, and certain reticular regions (4–7,16). Even after section behind the colliculi, a variety of spontaneous and reflex facial movements occur (10). However, there has been no unit-recording evidence identifying cells that discharge in relation to facial movement. In the present paper we report on the discovery of several cell types located within the medial pontomesencephalic reticular formation (RF) that discharge in relation to facial movements. We have previously presented preliminary data on a subgroup of these cells that discharge in association with pinna movement (15).

METHODS

Methods were as reported in the companion paper (14). Topical application of Xylocaine jelly, supplemented by subdermal injections of lidocaine, was used to block somatosensory input to receptive fields of facial cells. Event markers were employed to compare facial movement to polygraphically recorded electrooculogram, neck electromyogram, and unit discharge.

RESULTS

Cells related to movement of the facial musculature (n = 45) constituted 14.7% of the total number of cells encountered. Four distinct cell types were seen. These were: 1) cells related to movement of the ipsilateral eyelid, 2) cells related to movement of the ipsilateral lip region, 3) cells related to movement of the ipsilateral pinna, and 4) cells related to several facial movements. In comparison with other RF cells, a group, the facial movement cells were less likely to have tonic activity during sleep; i.e., type 2 cells were rare (P < 0.001, x²) and more likely to be active during the head-shake reflex (P < 0.01, x²). However, each cell subtype within the population of facial movement-related cells had quite distinctive behavioral correlates. Table 1 lists the frequency of cell subtypes and sleep and sensory responses.

response, and activity during head-shake reflex

<table>
<thead>
<tr>
<th>Sleep Type</th>
<th>% Total</th>
<th>n</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>Auditory</th>
<th>Shock</th>
<th>Strobe</th>
</tr>
</thead>
<tbody>
<tr>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Eyelid</td>
<td>2.2%</td>
<td>7</td>
<td>67*</td>
<td>0*</td>
<td>33*</td>
<td>0</td>
<td>0</td>
<td>53*</td>
</tr>
<tr>
<td>Lip</td>
<td>2.2%</td>
<td>7</td>
<td>57*</td>
<td>14*</td>
<td>29*</td>
<td>0</td>
<td>0</td>
<td>50</td>
</tr>
<tr>
<td>Pinna</td>
<td>7.1%</td>
<td>22</td>
<td>769</td>
<td>55</td>
<td>197</td>
<td>69*</td>
<td>891</td>
<td>781</td>
</tr>
<tr>
<td>Multiple</td>
<td>2.9%</td>
<td>9</td>
<td>133</td>
<td>88§</td>
<td>189</td>
<td>0</td>
<td>594</td>
<td>0</td>
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<tr>
<td>Total facial</td>
<td>14.7%</td>
<td>45</td>
<td>57</td>
<td>7</td>
<td>36</td>
<td>18</td>
<td>47</td>
<td>5</td>
</tr>
</tbody>
</table>

* and † Significant differences at 0.05 and 0.001 level between cell type and general RF cell population, and § Significant differences from other facial cell types at 0.05 and 0.001 level.
with the phasic relation of these cells to eye closure seen in waking. Eyelid movement cells were all located 1.2 ± 0.2 mm from the midline and between P3.5 and P6 in the gigantocellular tegmental field (FTG) (Fig. 2).

Ipsilateral lip movement cells

Seven cells (2.2% of the total number recorded) were related to movement of the ipsilateral lip. No cells showed any relation to movement of the contralateral lip. In five of the six cells the adequate movement was isolated to the upper lip or vibrissae, while in one, movement of both upper and lower lip was effective. As was the case with eyelid movement cells, unit discharge was related to the motor response rather than the stimulus. Manipulation of the vibrissae was ineffective unless it produced the appropriate movement of the lip. Stimulation of the skin of the lip with punctate pressure of as little as 150 mg produced a brisk response if the cat responded with movement of the ipsilateral lip. Conversely, even heavy pressure applied with the forefinger would not produce discharge if the appropriate contraction did not occur. Passive skin movement was ineffective. Injection of lidocaine and topical application of Xylocaine jelly to the lip and underlying tissues did not prevent unit discharge during movement in two units tested. Two cells were tested for response of 0.5-ms electric stimuli of sufficient intensity to cause twitching in the lip musculature. This stimulus did not produce unit discharge in either cell. As was the case with the eyelid cells, type 2 sleep patterns were rare in this group, while type 1 patterns were overrepresented (P < 0.02, x²). Auditory and visual responses were absent in all of these cells. Three of the six tested cells were active during execution of the head-shake reflex, not significantly different from the general cell population. All but one of the lip movement units, which were seen in five different cats, were grouped between P6 and P8 and between L1.2 and L2.3 (Fig. 2).

Ipsilateral pinna movement cells

A total of 22 cells (7.1% of the total recorded) discharged in conjunction with active movement of ipsilateral pinna. No cells discharging specifically in relation to contralateral pinna movement were seen. Unit activity in pinna movement cells was identical during both pinna movement elicited by auditory stimuli and movement elicited by irritating the hairs of the auricle with a cotton swab. Detailed observation of the topography of the pinna movement correlated with unit activity was made in 14 of these cells. Of these, four related to caudal, six to ventro-caudal, and four to rostral movements of the pinna. All but one of the cells related to pinna movement had little or no spontaneous activity when the pinna was in the normal intermediate "resting" position. Rapid active ear rotation in the appropriate direction was accompanied by a burst of unit discharge (Fig. 3). Five cells exhibited tonic discharge if the pinna was maintained by the cat in the displaced position but the remaining cells discharged only during the movement. Vigorous pinna movements were accompanied by bursts up to 30 spikes within a 0.5-s interval, while slower movements correlated with a single spike or no discharge at all. All of these cells were tested for response to somatic stimulation. When such stimulation elicited pinna movement, discharge occurred as in the case of spontaneous movements. In the absence of detectable ear movements, 16 of the 19 tested cells had no somatic field. Of the remaining cells, one responded to light manual stroking on any portion of the ipsilateral face. Two cells responded to blunt pressure applied to the area from the lower ipsilateral jaw to the region ventral to the pinna. These same three cells were totally unresponsive to punctate stimulation of up to 8 g applied to the same area with anesthesiometer. Two cells were tested for re-

FIG. 1. Response of eyelid movement cells to strobe stimulus. Each sweep has 100 ms duration.

FIG. 2. Anatomical distribution of facial cells. Circles, ipsilateral pinna movement cells; squares, ipsilateral lip movement cells; crossed squares, ipsilateral eyelid movement cells; triangles, nonspecific facial movement cells. Lines enclose concentrations of localized cell types. Lateralities of each cell type also differed (see text).

FIG. 3. Activity of pinna movement-related unit. Underline indicates when ipsilateral pinna was retracted to caudal position. Note the independence of unit activity from eye and head movements.
not change their base-line discharge rates during limb movements or movements of the caudal axial musculature.

The lack of facial movement specificity in these cells contrasted with the highly specific relations seen in all other facial cells. Not only did these cells discharge in conjunction with several movements, their relation to the movements was not directionally specific, as was the case in all other facial movement cells. For example, each of these cells accelerated during rostral, caudal, and ventral movements of the pinna.

This group of units also shared several other characteristics that distinguished them from other facial cells. Punctate stimulation of the ipsilateral face with as little as 100 mg produced unit discharge even if no movement was observed. However, the response was greatly enhanced if movement was evoked. Passive movement of either pinna in any direction produced tonic discharge similar to that occurring during active movement. This same manipulation was completely without effect in the ipsilateral pinna movement cells described above. Seven of the eight cells in this group responded to auditory stimuli, in contrast to the lack of response in other facial cells ($P < 0.001$, $x^2$). Six of the eight also responded to shock stimulation ($P < 0.05$, $x^2$). Although response latency was shortest to pulses applied to the ipsilateral facial field (1-2 ms), these cells also responded to stimulation of the torso and limbs, with an additional delay of 2-3 ms. Also in contrast to other facial cells was the lack of activity during the head-shake reflex in six of the seven cells tested ($P < 0.001$, $x^2$). Type 1 cells, which were overrepresented in the other facial cells, were significantly underrepresented in this group, while type 3 cells were overrepresented ($P < 0.001$, $x^2$). These cells were all clustered between P4 and P6.5 at 1.2 mm from the midline (Fig. 2).

**DISCUSSION**

Reticular facial movement-related cells were divisible into two distinct categories, those related to a single specific movement and those related to several movements. The range of motor relationships in the "specific" facial cells roughly corresponded to the range of movements known to be produced by the contraction of individual facial muscles. Pinna movements, which are controlled by a large proportion of the facial muscles, also comprise the largest portion of facial movement cells in the RF. Facial cells related to a single movement had few sensory responses, becoming active only when stimuli induced their specific motor correlate. The presence of activity during the reflex head shake can be explained by the facial and pinna movements consistently triggered during this reflex.

These cells were generally silent in rapid eye movement (REM) sleep, at a time when most surrounding cells achieve their highest activity levels (12). The spontaneous and reflex activity and lack of sensory response in these cells are similar to those seen in neighboring RF neurons related to eye movement (14). One may hypothesize that the functional role and synaptic mechanisms underlying the behavioral relations of the specific facial movement cells are analogous to the well-studied mechanisms of eye movement interneurons. Current anatomical evidence indicates the existence of synaptic connections from the RF regions containing facial cells to the facial nucleus (4, 5, 7, 16). Thus, some of these cells may relay their outputs over multisynaptic pathways.

Others have reported restricted facial receptive fields to "natural" stimuli in RF cells related to eye movements (1, 2, 9). These fields correspond in size to regions effective in eliciting movements in the present group of cells. However, such movements and spontaneous movements would not have been observable in these previous preparations. These investigations may therefore have been studying the sensory stimuli effective in triggering movements in the same cell population that we are reporting on here.

Cells related to several facial movements appear to form an anatomically and behaviorally distinct population from the specific facial movement cells. Unlike cells related to specific facial movements, they clearly receive a variety of exteroceptive and proprioceptive sensory inputs from the trigeminal, spinal, and auditory systems. The activity of these cells is qualitatively different from those of the specific movement cells in their lack of relation to the direction of movement as well as their relation to multiple movements. Their activity pattern is incompatible with any simple role in localized sensory or motor activity. Their discharge during spontaneous movement may be related to reflex discharge or facilitation of several groups of motoneurons. The presence of REM sleep activity and the absence of activity during vigorous head movement reflexes also distinguishes them from specific head movement-related cells.

Cells related to facial movement were distributed over a large proportion of the medial RF region studied, but there was a general localization of facial movement-related cells to RF regions rostral to the inferior olive (P8) and caudal to the level of the trochlear nucleus (P1). These cells were all rostral to the ipsilateral RF and largely medial to the facial nerve. Despite the large AP scatter of facial movement cells, the left-right segregation of subtypes was perfect. All the facial movement-related cells discharged exclusively or predominantly in relation to ipsilateral movements. The cells had different anatomical distributions within the ipsilateral RF. Cells related to pinna movement were most widely scattered. Pinna movement cells with virtually identical motor and sensory relations were separated by as much as 6 mm. In contrast, cells related to lip and eye movements formed fairly localized clusters. The cells having a nonspecific relation to a number of facial movements were also sharply localized to the region ventral to the abducens nucleus. This area has been shown to have nonsynaptic connections with a number of different motoneuron pools (8). Any understanding of the synaptic mechanisms mediating the observed behavioral relationships must await studies of the inputs and axonal trajectories of each cell type.

The present studies show the utility of investigating RF cells in unrestrained animals in a wide variety of movements. Several cell types appear to share certain sensory, sleep cycle, and movement characteristics. Cells sharing behavioral characteristics form clusters not otherwise apparent from histological examination of the RF. Our pre-